

Article

Not peer-reviewed version

A Unified Theory of Atomic and Molecular Structure

[Jiqing Zeng](#)* and Tianhe Zeng

Posted Date: 9 December 2025

doi: 10.20944/preprints202501.1033.v2

Keywords: electronic cloud model; hybrid orbital theory; molecular orbital theory; dynamic entity model of electron orbits; spatial configuration theory of electron orbits



Preprints.org is a free multidisciplinary platform providing preprint service that is dedicated to making early versions of research outputs permanently available and citable. Preprints posted at Preprints.org appear in Web of Science, Crossref, Google Scholar, Scilit, Europe PMC.

Copyright: This open access article is published under a [Creative Commons CC BY 4.0 license](#), which permit the free download, distribution, and reuse, provided that the author and preprint are cited in any reuse.

Disclaimer/Publisher's Note: The statements, opinions, and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions, or products referred to in the content.

Article

A Unified Theory of Atomic and Molecular Structure

Jiqing Zeng ^{1,*} and Tianhe Zeng ²

¹ South China Botanical Garden, Chinese Academy of Sciences, Guangzhou 510650, China

² College of Environment and Climate, Jinan University, Guangzhou 511443, China

* Corresponding author: zengjq@scib.ac.cn

Abstract

Revealing the structure of atoms and molecules has always been one of the important research goals in the field of quantum mechanics. The currently well-known atomic and molecular structure theories include Rutherford's planetary model, Bohr Sommerfeld atomic structure model, as well as atomic orbital theory, hybrid orbital theory, and molecular orbital theory. However, although these theories can explain atomic or molecular structures to some extent, they all have their own shortcomings, and there is currently no unified theory of atomic and molecular structures established. Here, we propose the Dynamic Entity Model of Electron Orbits, the Electron Spin Theory, and the Spatial Configuration Theory of Electron Orbits. Based on these new concepts and theories, we rearranged the extranuclear electrons of all elements in the periodic table, and explained the structure of atoms, the physical mechanisms of molecular formation, and the spatial structure of molecules. The theories of atomic and molecular structures based on quantum mechanics are often complex, difficult to understand, and inconsistent, while our new concepts and theories are based on classical physics and have the characteristics of being simple, intuitive, and easy to understand, and can logically and consistently explain the structure of atoms and molecules. Therefore, we have established a unified atomic and molecular structure theory based on the framework of classical physics, which has important scientific significance and application value.

Keywords: electronic cloud model; hybrid orbital theory; molecular orbital theory; dynamic entity model of electron orbits; spatial configuration theory of electron orbits

1. Introduction

In 1911, E. Rutherford proposed the "planetary model" of atomic structure, which suggested that electrons orbit the nucleus in the same way as planets orbit the sun¹. In 1913, Niels Bohr introduced the quantization hypothesis based on Rutherford's "planetary model" and proposed the Bohr model of hydrogen atomic structure². This model has achieved great success in explaining hydrogen atom spectra, but it is only applicable to hydrogen atoms and hydrogen like atoms, and cannot provide accurate explanations for more complex atomic structures. In 1916, Arnold Sommerfeld introduced the concept of elliptical orbits based on the Bohr model, making electron motion trajectories more realistic³. Sommerfeld's elliptical orbit theory can accurately calculate the energy levels of hydrogen atoms, highly consistent with experimental results, and successfully explain the fine structure in hydrogen atom spectra. Although Sommerfeld's theory presents difficulties in dealing with multi electron atoms, it can still provide predictions that are consistent with experimental results in certain specific situations. This fully demonstrates that Bohr and Sommerfeld's atomic structure models are quite close to physical reality.

However, Bohr and Sommerfeld's theories contain concepts such as quantization assumptions or conditions that cannot be explained by classical physics, resulting in their theories being in a semi classical and semi quantum state, which leads to contradictions and incompleteness in explaining microscopic phenomena. To address this contradiction and incompleteness, one approach is to explain quantized phenomena using the principles of classical physics; another is to abandon the

continuity theory of classical physics and establish a purely quantum mechanical theory. Max Planck, the founder of quantum mechanics, dedicated his entire life to explaining quantum phenomena using the continuity theory of classical physics, yet failed to achieve his goal. Instead, modern quantum mechanics, which rejects the continuity and determinism of classical mechanics, was quickly established. Quantum mechanics is considered a tremendous success in explaining atomic structure, as it can accurately predict and explain various experimental phenomena, such as atomic spectra and the fine structure of atomic energy levels. Consequently, Sommerfeld's theory was soon replaced by quantum mechanics and never continued to be refined and developed further.

Although the mathematical framework of quantum mechanics seems to demonstrate extremely high precision and accuracy in predicting and describing phenomena in the microscopic world, we often find ourselves in a dilemma when attempting to explain these complex phenomena using everyday language and common sense of classical physics. The interpretation of quantum mechanics has been a subject of extensive controversy and deep disagreements within the physics community. The popularity of the phrase "Shut up, and calculate!"⁴ in the quantum mechanics community fully reflects the profound confusion and difficulties encountered in interpreting quantum mechanics. This may suggest that the quantum mechanics theory centered on the Schrödinger equation is primarily a mathematical model rather than a purely physical one. Recently, our research has revealed that the diffraction-like and interference-like mechanisms of particle flows fundamentally refute the concept of wave-particle duality and De Broglie's hypothesis of matter waves, implying that the physical foundation upon which the Schrödinger equation is built does not hold⁵. This represents a revolutionary upset to quantum mechanics. Currently, there are numerous theories based on quantum mechanics for atomic and molecular structures, such as Valence Bond (VB) theory^{6,7}, Hybrid Orbital theory^{8,9}, Molecular Orbital (MO) theory^{10,11}, crystal field theory¹², and coordination field theory^{13,14}. While these theories appear prosperous on the surface, each has its own defects and shortcomings. Some theories even contradict one another, highlighting the imperfections of the current theoretical system and the lack of a unified theory for atomic and molecular structures. This means that the strategy of abandoning the continuity theory of classical physics to establish a purely quantum mechanics theory has ultimately proven unviable. Perhaps, we should revisit Planck's concept of energy quanta proposed in 1900, using classical physical principles to explain quantum phenomena and explore the correct path forward.

Recently, we have revised Planck's concept of energy quanta and the theory of radiation energy from accelerated motion of electrons^{15,16}. Based on classical physics principles, we have elucidated the physical mechanism of electron transitions and successfully derived and explained Bohr and Sommerfeld's atomic structure models. Our research shows that the motion of electrons outside the atomic nucleus is either a periodic circular (or elliptical) motion around the nucleus (steady state), or an accelerating or decelerating spiral motion (electron transition between steady states), all of which are continuous motions and do not have any discontinuity, discontinuity, jumping or uncertainty. The viewpoint on the intermittency, jumping, and uncertainty of electronic motion stems from the lack of a physical mechanism that can properly explain the phenomena of electronic transitions, orbital energy levels, orbital radii, and angular momentum quantization. We have successfully explained the quantization phenomena of hydrogen atoms and hydrogen-like atoms within the framework of classical physics, marking a fundamental solution to the semi-classical and semi-quantum problems in Bohr and Sommerfeld's theory. The core objective of this article is to construct a unified and comprehensive theory of atomic and molecular structures based entirely on the framework of classical physics.

2. Theoretical Framework

2.1. The Concept of Dynamic Entities and the Dynamic Entity Model of Electron Orbits

As is well known, material entities often have specific shapes and sizes, occupying a defined space. When an object undergoes high-speed periodic motion, the shape and size of the space

enclosed or occupied by its motion trajectory are also determined due to its fixed trajectory. For example, when a fan blade or helicopter propeller rotates at high speed, it appears to form a disk. Generally, when an object undergoes periodic motion, the space enclosed and defined by its motion trajectory is called the "periodic existence space" of the object. The periodic motion of an object includes vibration and rotation. When the periodic motion of an object is a reciprocating vibration, its periodic existence space is a linear structure (cylindrical structure). When the periodic motion of an object is circular or elliptical, its periodic existence space forms a circular ring or elliptical ring. When the circular ring or elliptical ring rotates around the diameter of the circle or the major axis of the ellipse, its periodic existence space becomes a sphere or ellipsoid. Therefore, when an object undergoes rapid periodic motion, its periodic existence space presents a specific shape and size. The dynamic distribution of an object in its periodic spatial existence is a "spatiotemporal whole", which we call a dynamic entity. The periodic existence space of objects can also be referred to as dynamic entity space. Obviously, dynamic entities are a special phenomenon formed by the high-speed periodic motion of objects. It can only be observed within the "time period" of the object's high-speed motion, whereas at a specific "instant" in time, only the moving object itself can be observed. Since the dynamic entity is a time-accumulated phenomenon resulting from the high-speed periodic motion of an object, its physical properties are the overall effects produced by this motion. The faster the periodic motion speed of an object, the shorter the motion period, and the more obvious the overall effect of the dynamic entity.

The circular or elliptical orbital motion of electrons around the nucleus is a typical high-speed periodic motion. The speed of electron movement around the nucleus is extremely fast (approximately $v_0=2.2\times 10^6$ m/s for electrons in the ground state of a hydrogen atom), while the radius of the electron orbit is extremely small ($r_0=0.53\times 10^{-10}$ m), and the motion period is extremely short ($T_0=6.6\times 10^{-17}$ s). Therefore, the electron orbit that moves in a circular or elliptical motion around the atomic nucleus can be regarded as a dynamic entity of a circular or elliptical ring.

Due to the susceptibility to electromagnetic fields generated by the motion of other electrons, the orbital plane of electrons in circular or elliptical motion around the nucleus is not always fixed and will rotate around the circular diameter or the long axis of the ellipse. Since the speed at which electrons move in a circular or elliptical motion around the atomic nucleus is much greater than the rotational speed of circular or elliptical orbits, it appears as if a dynamic entity of a circular or elliptical ring is rotating around its diameter or major axis, ultimately forming a spherical or ellipsoidal dynamic entity of electron motion orbit.

Assuming the period of circular or elliptical motion of electrons is T and the precession period is t , the charge passing through per unit time (charge flux) at the vertex of the long semi-axis of a sphere or ellipsoid is e/T , while the charge flux at other locations on the surface of the sphere or ellipsoid is $e/(T+t)$. Obviously, the charge flux per unit area (charge flux density) is highest at the vertex of the sphere or ellipsoid and decreases towards the middle of the sphere or ellipsoid. These characteristics of electronic orbital dynamic entities are of great significance.

Treating electron orbits as dynamic entities of spheres or ellipsoids greatly simplifies research and avoids complex and often unnecessary issues such as delving into the precise spatial position and momentum magnitude of electrons at any given moment. This new research approach allows us to focus more on the overall behavior and dynamic characteristics of electronic orbital dynamic entities, thereby more effectively revealing the essence of electronic motion. The concept of dynamic entities of electron orbits makes our understanding of the motion of electrons outside the atomic nucleus more intuitive and understandable. For example, when electrons absorb light quanta, the radius of their motion around the nucleus gradually increases, which can be understood as the dynamic entity of the electron orbit undergoing "expansion"; When electrons radiate light quanta, the radius of their motion around the nucleus gradually decreases, which can be understood as the dynamic entity of the electron orbit undergoing "contraction". The "expansion" and "contraction" of electronic orbital dynamic entities represent the absorption and radiation of energy, which is very similar to the thermal expansion and contraction phenomena of macroscopic objects.

The concept of dynamic entities is a physical model based on a holistic perspective, which provides us with a new perspective to examine the essence of macroscopic objects. Based on this model, it is easy to find that the periodic movement of high-speed microscopic particles forms dynamic entities, and countless microscopic dynamic entities aggregate to produce macro effects and form macro substances. The concept of dynamic entities builds a bridge connecting the micro and macro worlds, allowing us to gain a deeper understanding of the essence of the material world.

2.2. The Basic Physical Image of Atomic Structure

Based on the concept of dynamic entities and our previous research results, the basic physical image of atomic structure has been clearly presented:

Firstly, electrons move rapidly around the atomic nucleus, manifested as the nucleus being enveloped by a dynamic entity of spherical or ellipsoidal electron orbits, with the nucleus located at the center of the sphere or the focal point of the ellipsoid.

Secondly, in a multi electron atomic system, there are multiple electron orbital dynamic entities with the same electron layer outside the atomic nucleus, which inevitably repel each other. According to the principle of minimum energy, in order to minimize the energy of the atomic system, these electron orbital dynamic entities will be evenly and symmetrically distributed in space as much as possible.

Thirdly, in a multi electron atomic system, the dynamic entity of electron orbits is a layered structure resembling a Russian nesting doll (Figure 1).

Fourthly, the orbital radius of the first electron layer of element Z is r_1 , and the orbital radius of the n th electron layer is n^2r_1 . The electron orbital energy level of the first electron layer is E_1 , and the electron orbital energy level of the n th electron layer is E_1/n^2 .

$$r_1 = \frac{\epsilon_0 h^2}{\pi m Z e^2} \quad (1)$$

$$E_1 = -\frac{m e^4 Z^2}{8 \epsilon_0^2 h^2} \quad (2)$$

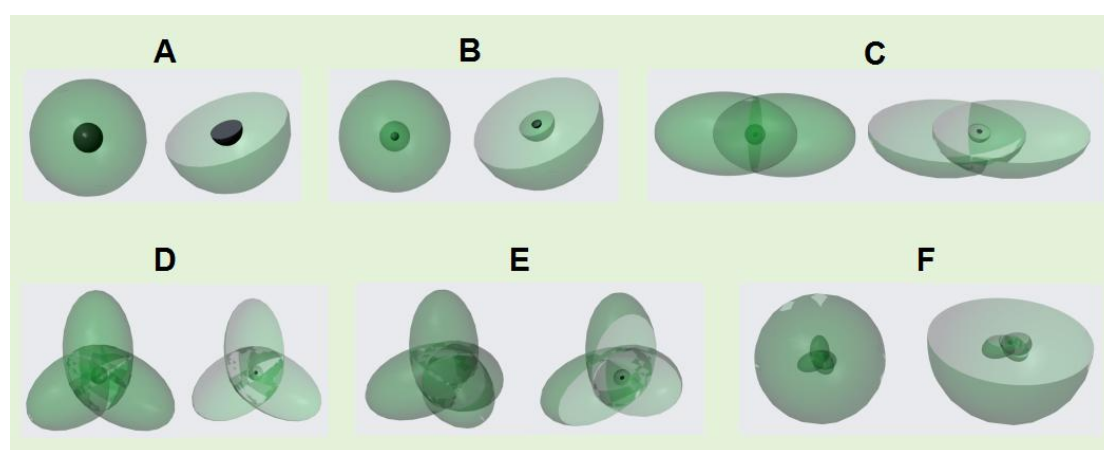


Figure 1. Dynamic entity image of atoms The dynamic entities and cross-sectional views of **A:** H and He atoms; **B:** Li and Be atoms; **C:** B and C atoms; **D:** N and O atoms; **E:** F and Ne atoms; **F:** Na and Mg atoms.

According to the principle of minimum energy and the distribution law of electron energy levels outside the atomic nucleus, the electron orbital energy level E_n in the n th electron layer cannot be lower than the electron orbital energy level E_1 in the first electron layer (i.e. $E_n \geq E_1$). Since the energy levels of electron orbits in the same electron layer are the same, the n th electron layer can accommodate a maximum of n^2 electron orbits ($n^2 E_n = E_1$) (since the energy levels of electron orbits are

negative, if the number of orbits is greater than n^2 , the energy level of the electron layer must be lower than E_1).

2.3. Electron Spin and Pairing

In the initial state, free electrons without force do not rotate or spin. When an electron moves around the nucleus, it rotates once as it moves around the nucleus, so the angular velocity ω of the electron's rotation is equal to the angular velocity ω of the electron's movement around the nucleus. This is the source of electron spin. The notion that electron spin is not rotation but an intrinsic property of electrons is clearly incorrect. Our latest research suggests that the correct concept of orbital magnetic moment should be the rotating electric quantity evr , the so-called Land factor (correction factor) g , and the spin quantum number $1/2$, which are erroneous conclusions based on the current erroneous concept of magnetic moment¹⁷.

According to the Existence Field Theory¹⁷, if an electron can serve as a stationary reference frame, then the interaction between electrons is called charge interaction (electrostatic force, Coulomb force). That is to say, Coulomb force only applies to a relatively stationary reference frame. If both electrons have spin motion, one of the spin electrons cannot be used as a stationary reference frame, so the interaction between the two spin electrons is not electrostatic force, but rather magnetic force generated by electron spin.

Electrons rotate synchronously around the atomic nucleus, generating a spin magnetic field. When two spin electrons are on the same electron orbit, they act like two small magnetic needles automatically reversing their magnetic poles and attracting each other to form an electron pair (Figure 2A). They move around the atomic nucleus once and rotate around each other once (Figure 2B). This is the physical mechanism of electron spin pairing on the same electron orbit.

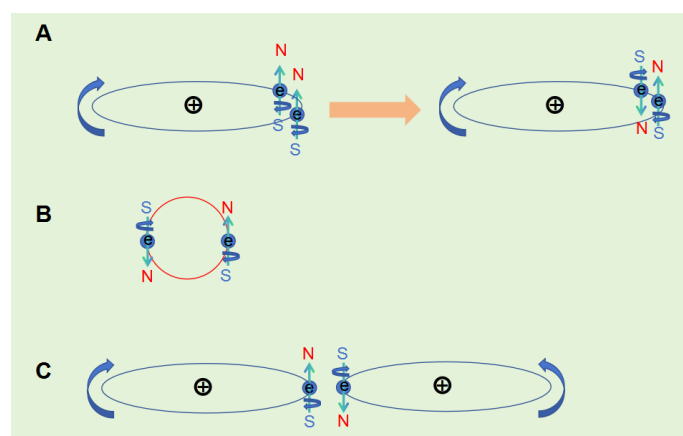


Figure 2. Electron spin and pairing. A: Electrons rotate synchronously around the atomic nucleus, generating a spin magnetic field. When two spin electrons are on the same electron orbit, they act like two small magnetic needles that automatically reverse their magnetic poles and attract each other to form an electron pair; B: Two spin electrons are attracted to each other by spin magnetic force, providing them with centripetal force to rotate around each other - when they move around the atomic nucleus together once, they also rotate around each other exactly once; C: A single electron orbital dynamic entity of two atoms is attracted to each other by the spin magnetic force of electrons, forming a covalent bond.

The faster the electron spin speed, the stronger the magnetic field generated, and the farther away from the spin electrons, the lower the magnetic field strength. According to our Existence Field Theory¹⁷, electron spin motion generates a "spin quantity" $Q_s = Q\omega$, which in turn generates a "spin quantity field". The strength of the spin quantity field is

$$E_s = k \frac{Q_s}{4\pi r^2} = k \frac{Q\omega}{4\pi r^2}$$

(3)

Among them, Q is the fundamental physical quantity (charge e or mass m), ω is the electron spin angular velocity, r is the radius of the sphere around the electron center, and k is the spin field constant. The interaction force between two spin quantities is

$$\mathbf{F} = \mathbf{Q}_s \times \mathbf{E}_s \quad (4)$$

The spin of charge generates charge spin quantity $Q_{se} = e\omega$, and the spin of mass generates mass spin quantity $Q_{sm} = m\omega$. Due to the fact that the interaction between charges is much greater than that between masses, we only consider the charge spin field (spin magnetic field) and its interaction. The strength of the electron spin magnetic field is:

$$\mathbf{E}_{se} = k_e \frac{e\omega}{4\pi r^2} \quad (5)$$

Among them, k_e is the spin magnetic field constant. The magnitude of the spin magnetic field interaction force (spin magnetic force) between two spin electrons is

$$F = Q_{se} E_{se} = k_e \frac{(\omega e)^2}{4\pi r^2} \quad (6)$$

When two electrons with opposite spins rotate around each other in a uniform circular motion under the action of spin magnetic force, the centripetal force is equal to the spin magnetic force F , and the circular motion radius is $r/2$, then:

$$F = k_e \frac{(\omega e)^2}{4\pi r^2} = m \frac{v^2}{r/2} \quad (7)$$

$$v = \omega e \sqrt{\frac{k_e}{8\pi m r}}$$

$$f = \frac{v}{2\pi r/2} = \frac{e\omega}{\pi} \sqrt{\frac{k_e}{8\pi m r^3}}$$

$$E_k = \frac{1}{2} m v^2 = \frac{1}{2} m \frac{(e\omega)^2 k_e}{8\pi m r} = \frac{1}{4} \times \frac{(e\omega)^2 k_e}{4\pi r}$$

$$E_p = -k_e \frac{(e\omega)^2}{4\pi r^2} \times \frac{r}{2} = -\frac{1}{2} \times \frac{(e\omega)^2 k_e}{4\pi r}$$

The total energy of a single spin electron is:

$$E = E_k + E_p = -\frac{1}{4} \times \frac{(e\omega)^2 k_e}{4\pi r} \quad (8)$$

Assuming two spin electrons are in a uniform circular motion around each other in the ground state orbit, with an energy of E'_0 , an orbital radius of $r'_0/2$, and a nuclear frequency of f'_0 . After absorbing energy ΔE within Δt time, electrons continuously increase their orbital radius and decrease their velocity. When the energy reaches its maximum value $E'_n=0$, its velocity is zero and its operating frequency $f'_n=0$. That is to say, when the electron reaches its maximum orbital, it is released from the spin magnetic force and the electron pair will separate. According to our previous research^{15,16}, we can conclude that:

$$\Delta E = E'_n - E'_0 = 0 - \left(-\frac{1}{4} \times \frac{(e\omega)^2 k_e}{4\pi r'_0}\right) = \frac{1}{2} \varepsilon f'_0 \Delta t \quad (9)$$

$$\frac{(e\omega)^2 k_e}{8\pi r'_0} = f'_0 (\varepsilon \Delta t)$$

$$f'_0 = \frac{\omega e}{\pi} \sqrt{\frac{k_e}{8\pi m r'_0{}^3}}$$

$$\frac{(e\omega)^2 k_e}{8\pi r'_0} = \frac{e\omega}{\pi} (\varepsilon \Delta t) \sqrt{\frac{k_e}{8\pi m r'_0{}^3}}$$

$$r'_0 = \frac{8(\varepsilon\Delta t)^2}{\pi m(e\omega)^2 k_e} \quad (10)$$

When $\Delta t=1s$, $\varepsilon\Delta t=\hbar$, then

$$r'_0 = \frac{8\hbar^2}{\pi m(e\omega)^2 k_e} \quad (11)$$

$$E'_0 = -\frac{1}{4} \times \frac{(e\omega)^2 k_e}{4\pi r'_0} = -\frac{m(e\omega)^4 k_e^2}{128\hbar^2} \quad (12)$$

We can imagine that spin electrons absorb light quanta within Δt time and then transition to higher energy levels E'_n or from higher energy levels to lower energy levels E'_m , releasing light quanta within Δt time:

$$\Delta E = E'_m - E'_n = -\frac{1}{4} \times \frac{(e\omega)^2 k_e}{4\pi r'_m} - \left(-\frac{1}{4} \times \frac{(e\omega)^2 k_e}{4\pi r'_n}\right) \quad (13)$$

$$\Delta E = \frac{(e\omega)^2 k_e}{16\pi r'_0} \left(\frac{1}{r'_n/r'_0} - \frac{1}{r'_m/r'_0}\right) \quad (14)$$

According to the electronic transition power $P = \frac{\Delta E}{\Delta t} = \varepsilon v$, let $\Delta t=1s$, $\varepsilon\Delta t=\hbar$, we can obtain:

$$v = Rc \frac{(\varepsilon_0 \omega^2 k_e)^2}{128} \left(\frac{1}{r'_n/r'_0} - \frac{1}{r'_m/r'_0}\right) \quad (15)$$

Among them, $R = \frac{me^4}{\varepsilon_0^2 \hbar^3 c}$ is the Rydberg constant. Let $R' = R \frac{(\varepsilon_0 \omega^2 k_e)^2}{128}$ be the Rydberg constant of the electron spin pairing system. Then

$$v = R' c \left(\frac{1}{r'_n/r'_0} - \frac{1}{r'_m/r'_0}\right) \quad (16)$$

Due to the atomic spectral structure conforming to the Rydberg formula, it can be obtained from equation (16)

$$r'_n = n^2 r'_0 \quad (17)$$

By substituting into equation (8), we can obtain

$$E'_n = \frac{E'_0}{n^2} \quad (18)$$

From equation (9), we can get

$$f'_n = \frac{f'_0}{n^3} \quad (19)$$

The angular momentum of spin electrons rotating around each other can be obtained by substituting the ground state orbital velocity v'_0 and orbital radius $r'_0/2$:

$$L'_0 = \frac{\hbar}{2\pi} = \hbar \quad (20)$$

According to equation (17), the same goes for:

$$L'_n = n \frac{\hbar}{2\pi} = n\hbar \quad (21)$$

It can be seen that two electrons with opposite spins rotate around each other under the action of spin magnetic force, just like the motion of electrons around the nucleus. Their energy levels, orbital radii, angular momentum, etc. are quantized, and the absorbed and radiated light quantum frequencies constitute the fine structure of atomic spectra. Based on the experimental data of the fine structure of atomic spectra, the corrected Rydberg constant R' can be calculated to determine the charge spin field constant k_e .

Due to the decrease in potential energy caused by spin magnetism between spin electrons, according to the principle of minimum energy, electrons outside the atomic nucleus will undergo spin pairing as much as possible to reduce the potential energy of the atomic system. Based on this,

it can also be inferred that, except for the outermost electron layer, the inner electron orbits must be paired double electron orbits, and the number of inner electrons must be even. In addition, based on the ground state orbital energy E'_0 of spin electrons rotating around each other, the electron pairing energy can be determined to be $2E'_0$.

When the number of electrons outside the atomic nucleus is odd, there must be a single-electron orbital in the ground state. When the atom is in an excited state, paired electrons will separate, forming two single-electron orbitals. According to the dynamic entity concept of electron orbits, the vertex positions (poles) of the spherical or ellipsoidal dynamic entity of a single-electron orbital possess the highest electron spin magnitude. Consequently, between the dynamic entities of single-electron orbitals from different atoms, the strongest spin magnetic force forms at their spherical or ellipsoidal vertex positions. Atoms with single-electron orbitals will thus bond through the spin magnetic force between their respective orbitals—this is the mechanism of electron spin pairing between different atoms (Figure 2C).

2.4. Space Configuration of Electron Orbits

Due to the mutual cancellation of magnetic moments between two electrons with opposite spin directions, the spin paired double electron orbits mainly exhibit charge characteristics. Therefore, there is only electrostatic repulsion between the third electron and the double electron orbits, and no spin magnetic force. It can be seen that there is electrostatic repulsion between double electron orbits or between double electron orbits and single electron orbits. Under the action of electrostatic repulsion, the electronic orbits will reach a balance of interaction forces, and their spatial distribution will eventually be in a uniform and symmetrical equilibrium state. This uniform symmetrical distribution state minimizes the energy of the entire atomic system.

The electronic orbital space configuration (abbreviated as "electronic configuration") is represented by V_{n-m} , where n is the number of electrons and m is the electronic configuration number. The larger the number of electrons n and the electronic configuration number m , the higher the potential energy of the electronic configuration.

1) 1 electronic

If there is only one electron in the outermost electron layer, it is obvious that there is only one electron orbital. The electronic orbit will adopt a uniformly symmetrical spherical or ellipsoidal orbital configuration. The electronic orbital configuration with only one electron on the electronic layer is represented by V_1 , the spherical orbital configuration is represented by V_{1-1} , and the ellipsoidal orbital configuration is represented by V_{1-2} .

2) 2 electronics

If there are two electrons in the outermost electron layer, it is easy to form electron spin pairing in the spin magnetic field, thereby reducing the orbital potential energy. Therefore, when the atom is in the ground state, the electron orbit is one spherical orbit or one ellipsoidal orbit, and the electron configurations are represented by V_{2-1} and V_{2-2} , respectively. When an atom is in an excited state, two spin paired electrons will separate, forming two single electron orbitals. There are two electronic configurations: spherical electron orbitals and ellipsoidal electron orbitals, as well as two symmetric distributions of ellipsoidal electron orbitals, represented by V_{2-3} and V_{2-4} , respectively.

3) 3 electronics

Due to the lowest energy of the system when atoms are in the ground state, electrons in the electron orbits are spin paired as much as possible, and the spatial distribution of electron orbits is as uniform and symmetrical as possible. Therefore, the three electrons in the outermost electron layer will form an electronic configuration of a spin paired spherical double electron orbit and an ellipsoidal single electron orbit (V_{3-1}), or an electronic configuration of a spin paired double electron ellipsoidal orbit and a linearly symmetrical distribution of a single electron ellipsoidal orbit (V_{3-2}). If the atom is in an excited state, two spin paired electrons will separate, and three electrons will form three single electron orbits, which can form one spherical orbital and two linearly symmetric

ellipsoidal orbits (V_{3-3}), or three coplanar ellipsoidal orbits with equal angles (120 degrees) between each pair of orbits in a uniform symmetric configuration (V_{3-4}).

4) 4 electrons

When an atom is in the ground state, the four electrons in the outermost electron layer can form two spin paired electron orbitals, including a spherical electron orbital and an ellipsoidal electron orbital (V_{4-1}), as well as two ellipsoidal electron orbitals symmetrically distributed (V_{4-2}). When the atom is in an excited state, the electronic configuration of the outermost electron layer varies depending on the energy of the excited state. When the energy of the excited state can only separate one electron pair, a spherical double electron orbital and two linearly symmetric ellipsoidal single electron orbital configurations (V_{4-3}) will be formed. When the energy of the excited state is more than enough to separate one electron pair but not enough to separate two electron pairs, three ellipsoidal electron orbits with uniform symmetry (V_{4-4}) will be formed (the angle between the electron orbits is equal to 120 degrees), including one double electron orbital and two single electron orbits. However, due to the higher charge of the double electron orbital compared to the single electron orbital, it will exert a squeezing effect on the two single electron orbits, reducing their angle (less than 120 degrees). When the energy of the excited state is sufficient to separate two electron pairs, four single electron orbits will be formed. According to the magnitude of the excitation energy, four single electron orbits will form one spherical orbital and three ellipsoidal orbital configurations (V_{4-5}) that are coplanar and have an orbital angle of 120 degrees, or four tetrahedral structures with equal orbital angles (109.47 degrees) (V_{4-6}), or two linearly symmetric ellipsoidal orbital configurations that are perpendicular to each other (V_{4-7}).

Figure 3 shows the electronic configuration of the outer nuclear electron layer with 1-8 electrons.

The electronic orbital spatial configuration is a reflection of the energy state of an atomic system. When an atom is in the ground state, the energy of the atomic system is at its lowest, electrons in the electron layer are spin paired as much as possible, the number of electron orbits in the valence layer is minimized, and the electron configuration is distributed as evenly and symmetrically as possible. When an atom is in an excited state, the energy of electrons outside the nucleus increases, causing the spin paired double electron orbits to split into two single electron orbits and increasing the number of valence electron orbits. As the number of electron orbits increases, the repulsive potential energy increases, and the spatial configuration of electron orbits changes from a low-energy configuration to a high-energy configuration. The changes in electron spin pairing energy levels and the transitions between different electron configurations are manifested in spectroscopy as fine and ultrafine structures of spectral lines.

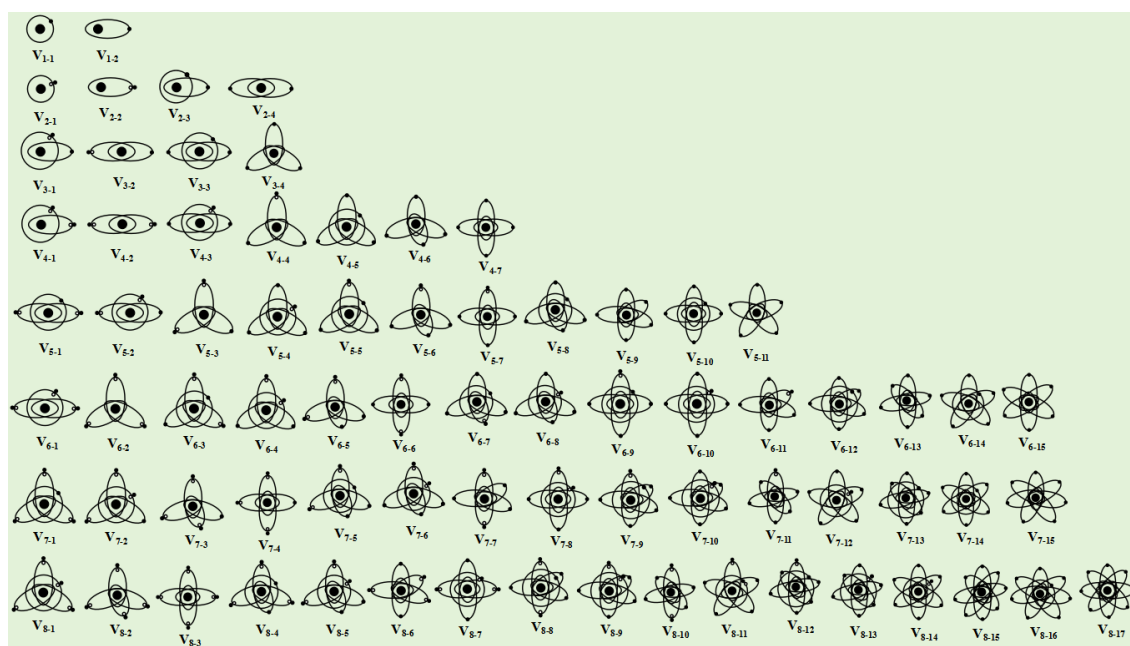


Figure 3. Valence electron orbital spatial configuration of atoms.

There are 1-8 electrons in the outer valence layer of the atomic nucleus, and their orbital spatial configurations are represented by V_{n-m} . Arranging from top to bottom, as the number of valence electrons n increases, the energy levels of the electron configuration also increase; From left to right, as the electronic configuration number m increases, the energy levels of the electronic configuration gradually increase.

2.5. Calculation of Atomic System Energy, Ionization Energy, and Electron Affinity Energy

From equation (2) and $E_n = E_1/n^2$, we can obtain the orbital energy levels of each electron layer of any atom. However, for multi electron atomic systems, the complex interactions between electrons make the calculation of the orbital energy levels of each electron layer more complicated. People usually use Slater's formula to calculate the energy of the electron layer⁹:

$$E_n^* = \frac{(Z-\sigma)^2}{n^2} R = \frac{Z^{*2}}{n^2} R \quad (22)$$

Among them, Z^* is the effective nuclear charge number, σ is the shielding coefficient, and $R = -13.6076 \text{ eV}$.

The so-called shielding effect refers to the shielding effect of inner layer electrons on outer layer electrons, for example, when the inner layer electrons are close to the atomic nucleus, they will cancel out the positive charge of the atomic nucleus. However, Slater's "shielding coefficient" σ not only includes the shielding effect of inner layer electrons, but also the interaction between electrons in the same electron layer. Obviously, this coefficient does not fully conform to the concept of shielding effect. In order to fully comply with the concept of shielding effect, we stipulate that the shielding coefficient σ only corresponds to the shielding effect of inner layer electrons on outer layer electrons, and no longer includes the interaction between electrons in the same electron layer. The interaction between electrons in the same electronic layer includes two aspects: one is the mutual repulsion between electron orbits, and the other is the mutual attraction between electrons with opposite spin directions. These interactions collectively shape the spatial configuration of electron orbits. Therefore, the electronic orbital energy level of the electronic layer should be composed of two parts, one is the electronic orbital energy level affected by the shielding effect, and the other is the electronic orbital spatial configuration potential formed by the interaction of electrons in the same electronic layer. The electronic orbital spatial configuration potential includes two parts: the electron orbital repulsion energy and the electron pairing energy.

Assuming that atom A of element Z has n electron layers, the n th electron layer has m electrons, and the electron pairing number is $\text{INT}(m/2)$. The orbital energy level of the n th electron layer of the ground state A atom is given by equation (22), the electronic spatial configuration potential energy of the n th electron layer is $E_{nV_m}(A)$, the electron orbital repulsion energy is E_{nRp} , and the electron pairing energy is $E_{np}(A)$. The total energy of the n th electron layer of atom A is:

$$E_n(A) = \frac{(Z-\sigma)^2}{n^2} R + E_{nV_m}(A) \quad (23)$$

$$E_{nV_m}(A) = E_{nRp} + \text{INT}(m/2)E_{np}(A) \quad (24)$$

The total energy of atomic system A is

$$E(A) = \sum_1^n E_i(A) \quad (25)$$

The average energy of each electron in the outermost electron layer (n th electron layer) is $E_n(A)/m$. Therefore, the first ionization energy of atom A is

$$I_{(1)}(A) = -E_n(A)/m \quad (26)$$

When losing k ($k < m$) electrons to form an A^{+k} ion, the average energy of each electron in the outermost electron layer is

$$E_n(A^{+k})/(m-k) = \frac{(Z-\sigma)^2}{n^2} R/(m-k) + E_{nV_{(m-k)}}(A)/(m-k) \quad (27)$$

The k th ionization energy is

$$I_{(k)}(A) = -E_n(A^{+k})/(m - k) \quad (28)$$

The m -th ionization energy is

$$I_{(m)}(A) = -E_n(A^{+m}) = -\frac{(Z-\sigma)^2}{n^2}R \quad (29)$$

The electron shielding coefficient σ within the n th electron layer is calculated using equation (29):

$$\sigma = Z - 0.2711n\sqrt{I_{(m)}(A)} \quad (33)$$

The energy of the first electron layer of atom A with atomic number Z greater than 2 is

$$E_1(A^{+Z-2}) = Z^2R + E_{1V_2}(A) = Z^2R + E_{1Rp}(A) + E_{1p} \quad (31)$$

Due to the completely uniform and symmetrical spatial configuration of a single spherical electron orbit, the electron orbit repulsion energy $E_{1Rp}(A) = 0$. The average energy of each electron is $E_1(A^{+Z-2})/2 = Z^2R/2 + E_{1p}/2$. According to the experimental data of ionization energy $I_{(Z-1)}$, the electron pairing energy of the first electron layer can be obtained as follows:

$$E_{1p}(A) = 2I_{(Z-1)} + Z^2R \quad (32)$$

When an atom has only one electron left, the shielding coefficient $\sigma = 0$, so the Z -th ionization energy is $I_{(Z)} = -Z^2R$.

Taking the 5th element (Boron, B) as a representative example, we perform the following calculations: The energy of the second electron shell in a boron atom is expressed as:

$$E_2(B) = \frac{(5 - \sigma)^2}{2^2}R + E_{2V_3}(B) = \frac{(5 - \sigma)^2}{2^2}R + E_{2Rp}(B) + E_{2p}(B)$$

$$E_2(B) = -3.4(5 - \sigma)^2 + E_{2V_3}(B) = -3.4(5 - \sigma)^2 + E_{2Rp}(B) + E_{2p}(B)$$

When only one electron remains in the second electron shell of the B atom, the energy of this electron shell becomes $-3.4(5 - \sigma)^2$. The energy required to remove this final electron (i.e., the third ionization energy) can be expressed as (36.93064eV) .

Using the experimentally measured third ionization energy of boron (36.93064eV), we derive the shielding constant σ for boron's inner electrons to be 1.70425. Thus,

$$E_2(B) = -36.93064 + E_{2V_3}(B)$$

So the average energy of each electron in the second electron layer is

$$\frac{E_2(B)}{3} = -12.3102 + \frac{E_{2V_3}(B)}{3}$$

According to the experimental data of the first ionization energy of atom B (8.29803eV), it can be concluded that

$$-12.3102 + E_{2V_3}(B)/3 = -8.29803$$

$$E_{2V_3}(B) = E_{2Rp}(B) + E_{2p}(B) = 12.0366\text{ (eV)}$$

When losing one electron to become a B^{+1} ion, the second electron layer has two electrons in a spherical two electron orbital, with an average energy of

$$E_2(B^{+1})/2 = -18.56532 + E_{2p}(B)/2$$

According to the experimental data of the second ionization energy of atom B (25.15484eV), it can be concluded that

$$E_{2p}(B) = -13.37904\text{eV}$$

By substituting into the previous equation ($E_{2Rp}(B) + E_{2p}(B) = 12.0366\text{eV}$), we obtain

$$E_{2Rp}(B) = 25.41564\text{eV}$$

When the B atom loses three electrons and becomes a B^{+3} ion, only two electrons from the first electron layer remain outside the nucleus, with an electron layer energy of

$$E_2(B^{+3}) = -5^2R + E_{1V1}(B) = -5^2R + E_{1p}(B)$$

$$E_2(B^{+3}) = -340.19 + E_{1p}(B)$$

The average energy per electron is

$$E_2(B^{+3})/2 = -170.095 + E_{1p}(B)/2$$

According to the experimental data of the 4th ionization energy of atom B (259.37521eV), it can be concluded that

$$E_{1p}(B) = -178.56024\text{eV}$$

Obviously, when the B atom loses 4 electrons and becomes a B^{+4} ion, only 1 electron from the first electron layer remains in the outer nucleus, with an energy of

$$E_2(B^{+4}) = -5^2R$$

$$E_2(B^{+4}) = -340.19$$

Therefore, the fifth ionization energy is

$$I_{(5)}(B) = -E_2(B^{+4}) = 340.19\text{eV}$$

As long as we obtain experimental data on the ionization energy of atoms, we can analyze the shielding coefficient of electrons in the outer and inner layers of the atomic nucleus, the potential energy of electron configurations in each layer, including the electron orbital repulsion energy and electron pairing energy, and obtain a clear understanding of the atomic structure. On the contrary, if theoretical parameters such as the shielding coefficient of electrons in the outer and inner layers of the atomic nucleus, the potential energy of electron configuration in the electron layer, etc. can be mastered, the ionization energy of atoms can also be predicted.

Based on the energy state of the atomic system, we can also calculate the electron affinity of elements. Assuming that atom A obtains an electron and becomes an A^{-1} ion, the energy of the n th electron layer is

$$E_n(A^{-1}) = \frac{(Z-\sigma)^2}{n^2}R + E_{nV_{(m+1)}}(A) = \frac{(Z-\sigma)^2}{n^2}R + E_{nRp}(A) + INT((m+1)/2)E_{np}(A)$$

(33)

Then, the first electron affinity is

$$E_{a(1)} = E_n(A^{-1}) - E_n(A) = \Delta E_V$$

(34)

$$\Delta E_V = E_{nV_{(m+1)}}(A) - E_{nV_m}(A) = \Delta E_{Rp} + \Delta E_p$$

(35)

$$\Delta E_{Rp} = E_{nRp_{(m+1)}}(A) - E_{nRp_m}(A)$$

(36)

$$\Delta E_p = INT((m+1)/2)E_{np}(A) - INT(m/2)E_{np}(A)$$

(37)

2.6. Atomic Spectrum

When the excited state of an atom only causes a change in the electronic configuration of the atomic valence layer:

The outermost electron number of atom A is m , the valence layer electron configuration is V_{m-i} , and the ground state electron configuration is V_{m-i} . When the atom absorbs energy and enters the excited state, its valence layer electron configuration changes from V_{m-i} to V_{m-j} . The energy level transitions from $E_{V_{m-i}}(A)$ to $E_{V_{m-j}}(A)$. The energy level difference is

$$\Delta E = E_{V_{m-j}}(A) - E_{V_{m-i}}(A) \quad (38)$$

The corresponding spectral frequency is

$$\nu = [E_{V_{m-j}}(A) - E_{V_{m-i}}(A)]/h \quad (39)$$

When the excitation of an atom causes a change in the radius and energy level (principal quantum number) of the electron orbit:

The total energy of the n th electron layer of atom A is

$$E_n(A) = \frac{(Z-\sigma)^2}{n^2} R + E_{nV_m}(A) + INT(m/2)E_{np}(A) \quad (40)$$

The valence layer electronic configuration energy of the n th electron layer is $E_{nV_{m-i}}(A)$, and the ground state valence layer electronic configuration energy is

$$E_{nV_{m-1}}(A) = E_{nV_m}(A) + INT(m/2)E_{np}(A) \quad (41)$$

The energy level of the valence layer electron configuration that transitions to the $(n+1)$ th electron layer is $E_{(n+1)V_{m-j}}(A)$, and the corresponding energy level change is

$$\Delta E = \frac{(Z-\sigma')^2}{(n+1)^2} R - \frac{(Z-\sigma)^2}{n^2} R + E_{(n+1)V_{m-j}}(A) - E_{nV_{m-i}}(A) \quad (42)$$

The corresponding spectral frequency is

$$\nu = [\frac{(Z-\sigma')^2}{(n+1)^2} R - \frac{(Z-\sigma)^2}{n^2} R + E_{(n+1)V_{m-j}}(A) - E_{nV_{m-i}}(A)]/h \quad (43)$$

2.7. Electron Arrangement Outside the Atomic Nucleus: Examples and Analysis

Based on the principle of minimum energy and the distribution law of electron energy levels outside the atomic nucleus, it is concluded that each electron shell can accommodate up to n^2 electron orbits, where n represents the shell number. Furthermore, according to the physical mechanism of electron spin pairing, each electron shell can hold a maximum of $2n^2$ electrons. The electron arrangement of atoms generally follows the principle of minimum energy, starting from the electron shell closest to the atomic nucleus and gradually progressing to higher energy levels. However, the specific number of electrons arranged in each electron shell does not directly correspond to the maximum electron capacity of each shell. As indicated by Equation (22), the energy gap between the first and second electron shells is relatively large, and as the shell number increases, the energy gap between adjacent electron shells diminishes. Therefore, when considering whether electrons should be arranged in the outermost or second outermost shell in higher electron shells, it is necessary to consider the energy of the entire atomic system. The most likely electron configuration is the one that minimizes the total energy of the atomic system.

For example, consider the element K with an atomic number of 19. The first electron layer can accommodate 2 electrons, the second electron layer can accommodate 8 electrons, and theoretically, the third electron layer can accommodate up to 18 electrons. Therefore, it seems that the third electron layer of K element can accommodate 9 electrons. But in reality, only 8 electrons are arranged in the third electron layer, and one electron is arranged in the fourth electron layer. This is because if there are 9 electrons arranged in the third layer, the potential energy of the electronic orbital spatial configuration ($E_{3V_9}(K)$) will be greater than the total potential energy of the electronic orbital spatial configuration of 8 electrons arranged in the third layer and 1 electron arranged in the fourth layer ($E_{3V_8}(K) + E_{4V_1}(K)$). In fact, the arrangement of 8 electrons in the third electron layer is the electron arrangement of the rare gas Ar. The atomic structure with the same arrangement of inner layer electrons as rare gas elements is called atomic core, and its valence electron orbital spatial configuration V_{8-1} has a perfectly uniform symmetrical structure and is very stable.

The electronic arrangement of transition elements, especially Group VIII elements, Group 1B elements, as well as lanthanide and actinide elements, is often highly controversial. The inner electron layers of these elements are not filled according to the maximum number of electrons that can be accommodated. As the atomic number increases, whether electrons are arranged in the unfilled inner electron layer or the outer electron layer needs to be determined based on the principle of the lowest energy in the atomic system. The increase in the number of inner electrons will increase the number of electron orbits, thereby enhancing the spatial configuration potential energy of electron orbits. Meanwhile, considering that electron spin pairing can reduce the potential energy of the system, the inner electron layer will be a spin paired double electron orbit. Therefore, as the atomic number increases, electrons will alternate between the inner electron layer or the outermost electron layer to keep the energy of the atomic system at its lowest state, until the inner electron layer forms an inert gas element atomic core structure and reaches a stable state. The electron arrangement of Group 1B elements is quite unique. Although the number of electrons outside the nucleus is sufficient to fill the 18 electrons in the second outer layer, this arrangement has a large potential energy for the electron orbital space configuration, and there is only one unpaired electron remaining in the outermost layer, which has a high energy and is therefore unstable. This type of electron arrangement with a non atomic core structure in the outer layer is unstable. Therefore, the 1B group elements continue the electron filling mode of the VIII group elements, forming a more stable atomic system with a secondary outer layer of 7 electron orbits (14 electrons) and an outermost layer of 5 electrons.

Based on the theory of electronic orbital spatial configuration, we have rearranged the extranuclear electron arrangement of all elements in the periodic table (Table 1), correcting the electron arrangement errors in the existing atomic orbital theory based on quantum mechanics. In the current electronic arrangement, some atoms exhibit an odd number of electrons in their inner shells, such as elements 39, 57, 58, 59, 65, 67, 69, 73, 75, and 77-89, 91-93, 95-97, 99, 101, and 103-112. Such arrangements contradict the principle of minimum energy because electron spin pairing reduces the potential energy of the atomic system, implying that the number of electrons in the inner shells must be even. Only after the inner shell electrons are spin-paired can unpaired electrons occupy the outermost shell.

To represent the electron arrangement outside the atomic nucleus, we can use the method shown in Figure 4 as needed. Figures 4a and b can be used to represent the number of electrons in each electron layer outside the atomic nucleus, while Figure 4c can be used to represent the spatial structure of the valence layer electron orbits of the atom.

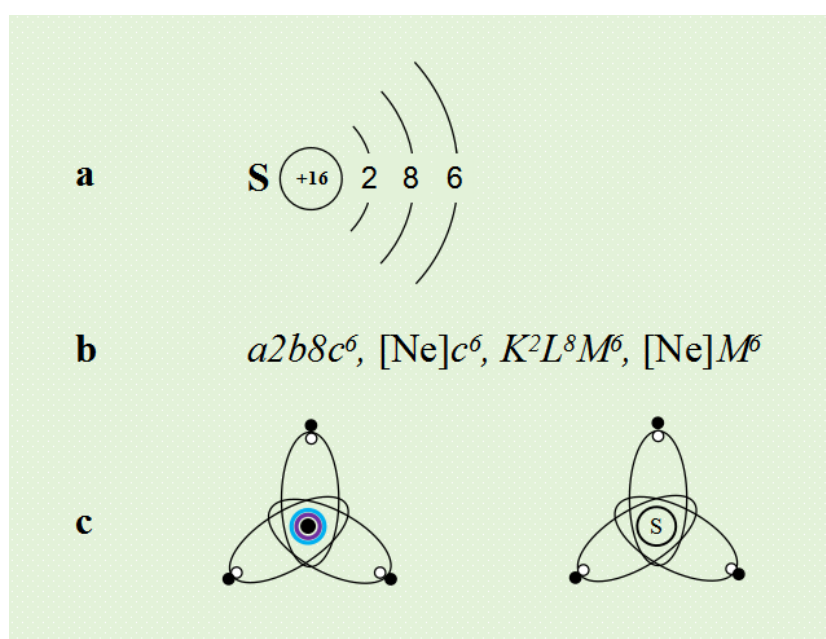


Figure 4. Representation of electron arrangement outside the atomic nucleus (taking sulfur atom S as an example) a, Atomic structure schematic diagram; b, Electronic layout: Use a, b, c, d, e, f, g or K, L, M, N, O, P, Q to represent 7 electron layers, and the number of electrons on each electron layer is represented by a number in the upper right corner of the electron layer symbol. If the inner layer is an atomic core structure, it can be abbreviated as [atomic core]; c, Valence electron orbital formula: The atomic nucleus is represented by a black circle, and the seven electron layers of a, b, c, d, e, f, g or K, L, M, N, O, P, Q are represented by circles in red, orange, yellow, green, cyan, blue, and purple colors, respectively; Alternatively, element names with circles can be used to represent the portion of the inner electron layer and atomic nucleus. Represent the outermost electron orbit with a circle or ellipse, and represent electrons with opposite spin directions with black and white dots.

Table 1. Exonuclear electron arrangement of each element in the periodic table.

Atomic Number	Element Symbol	Electron configuration	Atomic Number	Element	Electron configuration	Atomic Number	Element	Electron configuration
1	H	a^1	37	Rh	$a^2b^8c^{18}d^8e^1$	73	Ta	$a^2b^8c^{18}d^{32}e^8f^5$
2	He	a^2	38	Sr	$a^2b^8c^{18}d^8e^2$	74	W	$a^2b^8c^{18}d^{32}e^8f^6$
3	Li	a^2b^1	39	Y	$a^2b^8c^{18}d^8e^3$	75	Re	$a^2b^8c^{18}d^{32}e^8f^7$
4	Be	a^2b^2	40	Zr	$a^2b^8c^{18}d^8e^4$	76	Os	$a^2b^8c^{18}d^{32}e^8f^8$
5	B	a^2b^3	41	Nb	$a^2b^8c^{18}d^8e^5$	77	Ir	$a^2b^8c^{18}d^{32}e^{10}f^7$
6	C	a^2b^4	42	Mo	$a^2b^8c^{18}d^8e^6$	78	Pt	$a^2b^8c^{18}d^{32}e^{12}f^6$
7	N	a^2b^5	43	Tc	$a^2b^8c^{18}d^8e^7$	79	Au	$a^2b^8c^{18}d^{32}e^{14}f^5$
8	O	a^2b^6	44	Ru	$a^2b^8c^{18}d^8e^8$	80	Hg	$a^2b^8c^{18}d^{32}e^{18}f^2$
9	F	a^2b^7	45	Rh	$a^2b^8c^{18}d^{10}e^7$	81	Tl	$a^2b^8c^{18}d^{32}e^{18}f^3$
10	Ne	a^2b^8	46	Pd	$a^2b^8c^{18}d^{12}e^6$	82	Pb	$a^2b^8c^{18}d^{32}e^{18}f^4$
11	Na	$a^2b^8c^1$	47	Ag	$a^2b^8c^{18}d^{14}e^5$	83	Bi	$a^2b^8c^{18}d^{32}e^{18}f^5$
12	Mg	$a^2b^8c^2$	48	Cd	$a^2b^8c^{18}d^{18}e^2$	84	Po	$a^2b^8c^{18}d^{32}e^{18}f^6$
13	Al	$a^2b^8c^3$	49	In	$a^2b^8c^{18}d^{18}e^3$	85	At	$a^2b^8c^{18}d^{32}e^{18}f^7$
14	Si	$a^2b^8c^4$	50	Sn	$a^2b^8c^{18}d^{18}e^4$	86	Rn	$a^2b^8c^{18}d^{32}e^{18}f^8$
151617	P	$a^2b^8c^5$	51	Sb	$a^2b^8c^{18}d^{18}e^5$	87	Fr	$a^2b^8c^{18}d^{32}e^{18}f^8g^1$
18	S	$a^2b^8c^6$	52	Te	$a^2b^8c^{18}d^{18}e^6$	88	Ra	$a^2b^8c^{18}d^{32}e^{18}f^8g^2$
19	Cl	$a^2b^8c^7$	53	I	$a^2b^8c^{18}d^{18}e^7$	89	Ac	$a^2b^8c^{18}d^{32}e^{18}f^8g^3$
20	Ar	$a^2b^8c^8$	54	Xe	$a^2b^8c^{18}d^{18}e^8$	90	Th	$a^2b^8c^{18}d^{32}e^{18}f^8g^4$
21	K	$a^2b^8c^8d^1$	55	Cs	$a^2b^8c^{18}d^{18}e^8f^1$	91	Pa	$a^2b^8c^{18}d^{32}e^{18}f^8g^5$
22	Ca	$a^2b^8c^8d^2$	56	Ba	$a^2b^8c^{18}d^{18}e^8f^2$	92	U	$a^2b^8c^{18}d^{32}e^{18}f^8g^6$
23	Sc	$a^2b^8c^8d^3$	57	La	$a^2b^8c^{18}d^{18}e^8f^3$	93	Np	$a^2b^8c^{18}d^{32}e^{18}f^8g^7$
24	Ti	$a^2b^8c^8d^4$	58	Ce	$a^2b^8c^{18}d^{18}e^8f^4$	94	Pu	$a^2b^8c^{18}d^{32}e^{18}f^8g^8$
25	V	$a^2b^8c^8d^5$	59	Pr	$a^2b^8c^{18}d^{20}e^8f^3$	95	Am	$a^2b^8c^{18}d^{32}e^{20}f^8g^7$
26	Cr	$a^2b^8c^8d^6$	60	Nd	$a^2b^8c^{18}d^{20}e^8f^4$	96	Cm	$a^2b^8c^{18}d^{32}e^{22}f^8g^6$
27	Mn	$a^2b^8c^8d^7$	61	Pm	$a^2b^8c^{18}d^{22}e^8f^3$	97	Bk	$a^2b^8c^{18}d^{32}e^{24}f^8g^5$
28	Fe	$a^2b^8c^8d^8$	62	Sm	$a^2b^8c^{18}d^{22}e^8f^4$	98	Cf	$a^2b^8c^{18}d^{32}e^{26}f^8g^4$
29	Co	$a^2b^8c^{10}d^7$	63	Eu	$a^2b^8c^{18}d^{24}e^8f^3$	99	Es	$a^2b^8c^{18}d^{32}e^{28}f^8g^3$
30	Ni	$a^2b^8c^{12}d^6$	64	Gd	$a^2b^8c^{18}d^{24}e^8f^4$	100	Fm	$a^2b^8c^{18}d^{32}e^{28}f^8g^4$

31	Cu	$a^2b^8c^{14}d^5$	65	Tb	$a^2b^8c^{18}d^{26}e^8f^3$	101	Md	$a^2b^8c^{18}d^{32}e^{30}f^8g^3$
32	Zn	$a^2b^8c^{18}d^2$	66	Dy	$a^2b^8c^{18}d^{26}e^8f^4$	102	No	$a^2b^8c^{18}d^{32}e^{30}f^8g^4$
33	Ga	$a^2b^8c^{18}d^3$	67	Ho	$a^2b^8c^{18}d^{28}e^8f^3$	103	Lr	$a^2b^8c^{18}d^{32}e^{32}f^8g^3$
34	Ge	$a^2b^8c^{18}d^4$	68	Er	$a^2b^8c^{18}d^{28}e^8f^4$	104	Rf	$a^2b^8c^{18}d^{32}e^{32}f^8g^4$
35	As	$a^2b^8c^{18}d^5$	69	Tm	$a^2b^8c^{18}d^{30}e^8f^3$	105	Db	$a^2b^8c^{18}d^{32}e^{32}f^8g^5$
36	Se	$a^2b^8c^{18}d^6$	70	Yb	$a^2b^8c^{18}d^{30}e^8f^4$	106	Sg	$a^2b^8c^{18}d^{32}e^{32}f^8g^6$
	Br	$a^2b^8c^{18}d^7$	71	Lu	$a^2b^8c^{18}d^{32}e^8f^3$	107	Bh	$a^2b^8c^{18}d^{32}e^{32}f^8g^7$
	Kr	$a^2b^8c^{18}d^8$	72	Hf	$a^2b^8c^{18}d^{32}e^8f^4$	108	Hs	$a^2b^8c^{18}d^{32}e^{32}f^8g^8$

3. Theoretical Application and Verification

3.1. The Formation and Essence of Chemical Bonds

Molecules are structural forms formed by the combination of atoms. According to the principle of minimum energy, atoms can combine with each other to form stable molecules, and the energy (E_M) of the molecular system must be less than the sum of the energies $E_{A(i)}$ of each atomic system before the formation of the molecule:

$$(44) \quad E_M < \sum_1^n E_{A(i)}$$

If the energy of a molecule composed of atoms combined with each other is less than the total energy of the ground states of each atom, it must be due to gravitational interactions between the atoms, thereby reducing the potential energy of the system. Obviously, the gravitational interaction between atoms must be electromagnetic interaction, either electrostatic attraction or magnetic attraction.

Generally, the outermost electrons of metal elements have higher energy levels. When the atomic system is in a higher energy state, these electrons are prone to becoming free electrons, leaving the metal atoms as positively charged cations. If a large number of identical excited-state metal atoms gather together, during the de-excitation process, the metal cations and free electrons will bind together through electrostatic attraction, forming metal elemental molecules. If there are different metal atoms in the atomic system, a metal alloy is formed. The electrostatic attraction between metal cations and free electrons is called a metal bond.

Usually, the outermost electron energy of non-metallic elements is low. When there is a single electron orbital in their ground state valence layer electron configuration, obtaining one electron can spin pair into a double electron orbital, thereby reducing the potential energy of the atomic system and forming a more stable anion. For instance, halogen elements like the chlorine atom (Cl) have 7 electrons in their outermost electron shell, with a valence electron configuration of V_{7-1} or V_{7-2} . There are 4 electron orbits in its ground state, among which 3 are paired electron orbits and 1 is a single electron orbital. If the chlorine atom gains 1 electron, its valence electron configuration will change to V_{8-1} , resulting in 4 paired electron orbits and a further reduction in potential energy. This transforms it into a stable [Ar] atomic core structure, forming a negatively charged chloride ion (Cl^-). The outermost electrons of metal atoms has higher electron energy, especially for alkali metal atoms such as sodium (Na), which has only one electron in its outermost shell, with the inner electron configuration forming a stable [Ne] atomic core structure. Consequently, the outermost electron is highly susceptible to becoming a free electron, forming a positively charged sodium ion (Na^+). When metal cations (Na^+) and non-metal anions (Cl^-) are mixed together, they attract each other through electrostatic forces, further lowering their potential energy and thereby forming a stable ionic compound molecule ($NaCl$) (Figure 5). Positive and negative ions are attracted to each other by electrostatic forces. When the positive and negative ions approach each other, electrostatic repulsion is generated due to the negative charge of the electron orbital dynamic entity. When the electrostatic

attraction between positive and negative ions is balanced with the electrostatic repulsion between the electron orbital dynamic entity, a stable ionic bond is formed.

If the outer electrons of an atom do not become free electrons to form cations, nor gain electrons to become anions, the atom remains electrically neutral, and thus atoms cannot attract each other through electrostatic forces. Obviously, the binding force between electrically neutral atoms can only be magnetic attraction. Since the magnetic moment of an atom primarily arises from the electron spin in the single-electron orbits of the outermost electron shell, when atoms with single electron orbits in the valence electron configuration approach each other, the dynamic entities of single electron orbits between atoms will be attracted to each other by spin magnetic force. When the spin magnetic force and electrostatic repulsion between single electron orbital dynamic entities reach equilibrium, stable **covalent bonds** are formed.

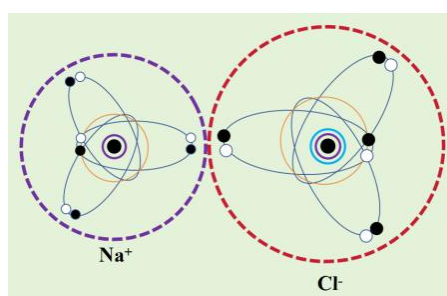


Figure 5. Formation of Ionic Bonds: Taking Sodium Chloride Molecules as an Example.

The rotational motion of atoms or ions causes their dynamic entities to take on a spherical shape. Sodium ions (Na^+) carry a positive charge as a whole, represented by a purple dashed circle; The chloride ion (Cl^-) carries a negative charge as a whole, represented by a red dashed circle. Sodium ions and chloride ions approach each other through electrostatic attraction, but they cannot approach infinitely because the two electron orbits between sodium ions and chloride ions are dynamic entities with negative charges, and there is electrostatic repulsion between them. When the electrostatic attraction and electrostatic repulsion between sodium ions and chloride ions reach an equilibrium state, stable ionic bonds are formed.

3.2. Spatial Structure of Molecules

3.2.1. Molecular Structures of Ionic Crystals and Metal Crystals

Under the electrostatic attraction between cations and anions, a structure with a uniform and symmetrical distribution of cations and anions is formed, which is known as the ionic crystal structure. This structure minimizes the electrostatic potential energy between cations and anions within the ionic crystal. The structure of an ionic crystal depends on the amount of charge carried by the ions and the radii of the cations and anions. The charge of the ions determines the strength of the electrostatic interaction between them, while the ion radii influence the packing arrangement of ions in the crystal. In addition, the shape of ions, i.e. the electronic orbital spatial configuration, also affects their arrangement in crystals, forming specific geometric structures (Figure 6a, 6b).

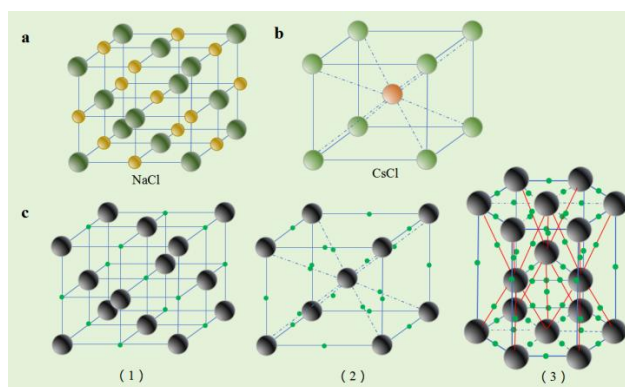


Figure 6. Ionic Crystal and Metal Crystal Structures. A: NaCl crystal structure; B: CsCl crystal structure; C: Metal crystal structure, (1) metal crystal structure composed of +1 valent cation and one free electron, (2) metal crystal structure composed of +2 valent cation and one free electron, (3) metal crystal structure composed of +3 valent cation and one free electron.

Similar to the structure of ionic crystals, a large number of metal cations and free electrons tightly bind together under electrostatic attraction to form a regularly arranged crystalline structure. Both in metal crystals and ionic crystals, the arrangement of atoms follows the principle of minimum energy, forming a uniform and symmetrical ordered crystalline structure through closest packing, thereby reducing the potential energy of the entire system. It is commonly believed that free electrons in metal crystals can move freely, randomly, and disorderly between metal atoms. In fact, the "free electrons" in a metal lattice are considered "free" only relative to the atomic nuclei of the metal atoms, that is, they have broken free from the atomic nuclei's restraint and no longer orbit around the atoms, but they are still bound to the metal cation through electrostatic attraction. This means that these electrons are not completely free to move randomly within the metal lattice but instead vibrate within a certain range. In ionic crystals, anions and cations are bound to each other by electrostatic attraction, while in metal crystals, metal cations repel each other. Therefore, free electrons are necessary to provide electrostatic attraction between metal cations to bind them. A metal crystal composed of a +1 valent cation and one free electron, with 4 free electrons evenly and symmetrically distributed around the metal cation (Figure 6c (1)). This is similar to the structure of NaCl type ionic crystals (Figure 6a), except that in the metal crystal, free electrons replace anions. A metal crystal composed of a +2 valent cation and one free electron, with 8 free electrons evenly and symmetrically distributed in space around the metal cation (Figure 6c (2)). A metal crystal composed of a +3 valent cation and one free electron, with 16 free electrons evenly and symmetrically distributed in space around the metal cation (Figure 6c (3)).

3.2.2. The Molecular Structure of Covalent Compounds

Due to the highest charge flux at the vertex of the ellipsoid of a single electron orbital dynamic entity, the electron spin field density per unit time is highest at that location. If the ellipsoidal single electron orbital dynamic entity of another atom approaches, then the two ellipsoidal single electron orbital dynamic entities must have the maximum **spin magnetic force** at the vertex in a "head to head" manner, which determines the directionality of the covalent bond. When single electron orbits are attracted to each other by spin magnetic force, if the symmetry axes of the two single electron orbits are connected to the atomic nucleus on the same straight line, the covalent bond formed in this way is called an α bond (Figure 7a). If the symmetry axes of the two single electron orbits are not connected to the atomic nucleus on the same straight line, the covalent bond formed in this way is called a β bond (Figure 7b).

Assuming that A atom forms an α bond with B atom, and the "charge spin quantity" of A atom's single electron orbit is $Q_{S(A)} = e\omega_A$, and the "charge spin quantity" of B atom's single electron orbit is $Q_{S(B)} = e\omega_B$, then the spin magnetic force of their spin pairing is:

$$F = k_e \frac{e^2 \omega_A \omega_B}{4\pi r^2} \quad (45)$$

Due to the coincidence of the electron orbital symmetry axis of the α bond with the line connecting the two atomic nuclei of AB, the spin magnetic force of spin pairing is the binding force between AB atoms.

Assuming that A atom and B atom form a β bond, and the angle between the symmetry axis of the single electron orbit and the line connecting the atomic nucleus is θ , the binding force between AB atoms is $F \cos\theta$. Therefore, the beta bond is slightly weaker than the α bond.

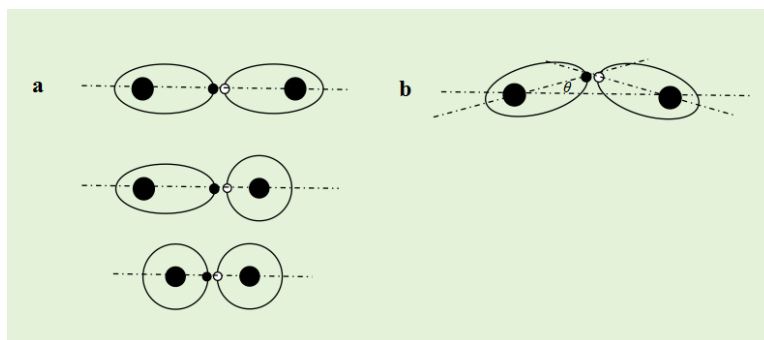


Figure 7. Spin pairing mode of single electron orbits between atoms **a**, The symmetry axes of two single electron orbits are aligned with the line connecting the atomic nucleus. The bond formed by this spin pairing method is called an α bond. **b**, The symmetry axes of two single electron orbits are not on the same straight line as the line connecting the atomic nucleus. The bond formed by this spin pairing method is called a β bond.

Based on the sum of the energies of each atomic system and the difference in energy between the molecular system, we can determine the bond energy of covalent bonds

$$E_c = \sum_1^n E_{A(i)} - E_M \quad (46)$$

If an atom has only one single electron orbital in its valence electron configuration, then the covalent bond it forms with other atom that has only one single electron orbital must be an α bond. For example, a hydrogen atom has only one single electron orbital, and any covalent bond formed between it and any atom is an α bond. In molecules such as SF₆, BF₃, and PCl₃, the valence layer electron configuration of F and Cl atoms is V₇₋₂, with only one single electron ellipsoidal orbital. Therefore, the S-F, B-F, and P-Cl covalent bonds are all α bonds.

If an atom has multiple single electron orbits in its valence layer, then the covalent bond it forms with other atoms that only have one single electron orbital must also be an α bond. For example, the valence layer electron configuration of carbon atoms in methane molecules (CH₄) is V₄₋₆, with four ellipsoidal single electron orbits that spin pair with the single electron orbits of four hydrogen atoms to form four covalent bonds, all of which are α bonds.

Since an atom with multiple single electron orbits can form covalent bonds with multiple atoms, such an atom is the central atom of covalent molecule. The spatial configuration of the valence electron orbits of the central atom usually determines the spatial structure of the molecule. For example, the electronic orbital spatial configuration of the carbon atom valence layer in methane molecules is V₄₋₆, which is a tetrahedral spatial structure composed of four ellipsoidal single electron orbital dynamic entities. Therefore, the spatial configuration of methane molecules is tetrahedral. The electronic configuration of the valence layer of sulfur atoms in SF₆ molecules is V₆₋₁₂, which is a regular octahedral structure composed of six single electron orbits. Therefore, the spatial structure of SF₆ molecules is a regular octahedron. The valence layer electronic configuration of boron atoms in BF₃ molecule is V₃₋₄, which is an equilateral triangle structure composed of three ellipsoidal single electron orbital dynamic entities. Therefore, the spatial structure of BF₃ molecule is a planar equilateral triangle. The valence layer electron configuration of phosphorus atom in PCl₅ molecule is V₅₋₉, which

is a triangular bipyramid structure composed of five single electron orbits. Therefore, the spatial structure of PCl_5 molecule is a triangular bipyramid (Figure 8).

The valence layer electron configuration of nitrogen atoms in ammonia molecules (NH_3) is $V_{5,6}$, which is a tetrahedral structure composed of three single electron orbitals and one double electron orbital. Due to the fact that the double electron orbitals do not participate in the formation of covalent bonds, the spatial structure of ammonia molecules is not a regular tetrahedron but a triangular pyramid. Due to the higher charge of the double electron orbital dynamic entity compared to the single electron orbital, it forms a stronger electrostatic repulsion squeezing effect on the other three single electron orbitals. Therefore, the H-N-H bond angle is not 109.47° , but 107° . Similarly, the valence layer electronic configuration $V_{5,6}$ of phosphorus atoms in phosphine molecules (PH_3) is a tetrahedral spatial structure composed of three ellipsoidal single electron orbitals and one ellipsoidal single electron orbital dynamic entity. Due to the absence of double electron orbitals in the formation of covalent bonds, the spatial structure of PH_3 molecule is not a regular tetrahedron but a triangular pyramid, and the H-P-H bond angle is not 109.47° , but 93.5° . The valence layer electron configuration of oxygen atoms in water molecules (H_2O) is $V_{6,5}$, which is a tetrahedral structure composed of two single electron orbitals and two double electron orbitals. Due to the fact that two double electron orbitals do not participate in the formation of covalent bonds, only two single electron orbitals form covalent bonds, the spatial structure of water molecules is a triangular structure. Due to the electrostatic squeezing effect of the two electron orbital dynamic entity, the H-O-H bond angle of water molecules is not 109.47° , but 104.5° (Figure 8).

If an atom has multiple single electron orbits in its valence electron configuration, it may form 2 or 3 covalent bonds when it approaches another atom that also has multiple single electron orbits. For example, the electronic configuration of the valence layer of an excited oxygen atom is $V_{6,3}$, with 2 double electron orbits and 2 single electron orbits. Two oxygen atoms form two α bonds by spin pairing with an ellipsoidal single electron orbit and a spherical single electron orbit, respectively, and combine to form an oxygen molecule (O_2) (Figure 8). Due to the exposure of spherical single electron orbits in the outer layer of the molecule, oxygen molecules are susceptible to the influence of external magnetic fields, thus exhibiting paramagnetism. In addition, due to the large distance between the spherical single electron orbits of two oxygen atoms, the formed α bond is weak and easy to break, thus oxygen molecules have strong chemical activity. When two α bonds are formed between atoms due to the simultaneous existence of ellipsoidal and spherical single electron orbits, we call the α bond formed between ellipsoidal single electron orbits "type I α bond" and the α bond formed between spherical single electron orbits "type II α bond".

If there are multiple ellipsoidal single electron orbits in the valence layer electronic configuration of an atom, due to the presence of certain angles in their spatial distribution, the 2 or 3 covalent bonds formed between such atoms must be β bonds. For example, the valence layer electron configuration of an excited nitrogen N atom is $V_{5,6}$, with one ellipsoidal double electron orbits and three ellipsoidal single electron orbits. Two nitrogen atoms spin pair with three ellipsoidal single electron orbits to form three β bonds, forming a nitrogen molecule (N_2). Nitric oxide molecule (NO) forms two β bonds between the two ellipsoidal single electron orbits of N atom and the two ellipsoidal single electron orbits of O atom. Due to the presence of an unpaired single electron orbital in the N atom, NO molecules exhibit paramagnetism. The valence layer electronic configuration of carbon atoms in carbon monoxide (CO) molecules is $V_{4,4}$, and the valence layer electronic configuration of oxygen atoms is $V_{6,4}$. The two ellipsoidal single electron orbits of C atoms and the two ellipsoidal single electron orbits of O atoms form two β bonds. There are no unpaired single electron orbits in CO molecules, therefore it has diamagnetism. In carbon dioxide molecules (CO_2), the valence layer electron configuration of carbon atoms is $V_{4,6}$, with 4 single electron orbits. One carbon atom can form 4 β bonds with 2 oxygen atoms, and the O-C-O bond angle is 180° . The spatial structure is linear. The valence layer electronic configuration of sulfur atoms in sulfur trioxide molecules (SO_3) is $V_{6,14}$, with six single electron orbits uniformly distributed in the same plane at an angle of 60° . Therefore, one

sulfur atom can form six β bonds with three oxygen atoms, with an O-S-O bond angle of 120° . The spatial structure of SO_3 molecules is a planar triangle (Figure 8).

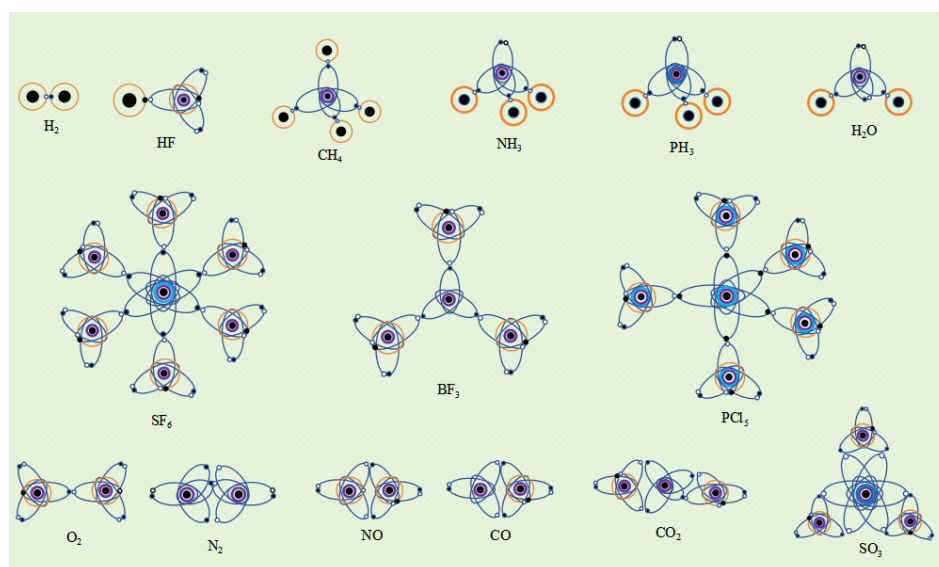


Figure 8. Structures of Several typical Covalent Molecules (Electronic Orbital Formulas).

The two carbon atoms of ethylene molecule (C_2H_4) have a V_{4-5} valence layer electronic configuration. The two ellipsoidal single electron orbits in the valence layer of each carbon atom spin pair with the single electron orbits of hydrogen atoms to form two "C-H" type I α bonds. Two carbon atoms form a "C-C" type I α bond with ellipsoidal single electron orbit spin pairing, and a "C-C" type II α bond with spherical single electron orbit spin pairing. Type II α bonds are weaker than type I α bonds and are more prone to breakage, making ethylene molecules more reactive. The two carbon atoms of acetylene molecule are also in the V_{4-5} valence layer electronic configuration, forming two "C-C" β bonds and one "C-C" α bond between the two carbon atoms. Due to the presence of more β bonds in the covalent bonds of acetylene molecules, they are more reactive than ethylene molecules. The carbon atoms in butadiene (including trans and cis 1,3-butadiene) are also in the V_{4-5} valence layer electronic configuration. In addition to forming an I type α bond, adjacent carbon atoms also form a "C-C" II type α bond. The four carbon atoms in the butadiene molecule are tightly bound together by three type I α bonds and three type II α bonds, further reducing the energy of the molecular system. Therefore, the butadiene molecule has high stability. The valence layer electronic configuration of the six carbon atoms in benzene molecule (C_6H_6) is V_{4-5} , with each carbon atom having a "C-C" type I α bond and a "C-C" type II α bond with adjacent carbon atoms. In addition, the spherical single electron orbits between carbon atoms in opposite positions on the benzene ring have opposite electron spin directions, resulting in spin magnetism and the formation of "C-C" type II α bonds. In this way, a strong bond is formed between the six carbon atoms of the benzene ring molecule, and the potential energy of the molecular system is greatly reduced, making the benzene ring molecule very stable (Figure 9).

The electronic configuration of the valence layer of an atom reflects its energy state. Different energy states determine the different valence electron configurations of atoms, and when covalent compound molecules are formed, they form different spatial structures. Therefore, the spatial structure of a molecule is essentially determined by the energy state of the atoms at the time of its formation. For example, in excited states with lower energy levels, the valence electron configuration of carbon atoms is V_{4-5} , and adjacent carbon atoms can form type I α bonds and type II α bonds. Six carbon atoms can form a stable hexagonal structure like a benzene ring. By using this as a unit, a planar network structure composed of regular hexagonal structures can be formed, which is the basic structure of graphene molecules. Graphene with a planar network structure is stacked layer by layer, and the spherical single electron orbits of carbon atoms between layers exhibit spin magnetic

interactions, forming sheet-like graphite. If the edges of such a planar network structure are raised and connected end-to-end, fullerene molecular structures can be formed, which can be spherical, ellipsoidal, cylindrical, or tubular in shape. At higher energy levels of excitation, the valence electron configuration of carbon atoms changes to V_{4-6} , with each carbon atom serving as the central atom and forming four Type I α bonds with four carbon atoms. The four carbon atoms form a three-dimensional network structure at the vertices of a regular tetrahedron, which is known as the diamond molecular structure (Figure 9). Due to the strong covalent bonds formed between each carbon atom and four carbon atoms, releasing a large amount of energy, the spatial potential energy of the entire molecular system is extremely low, making diamond molecules very stable. Elemental allotropes like graphene and diamond are giant covalent molecules with different structures formed under excited states of different energy states.

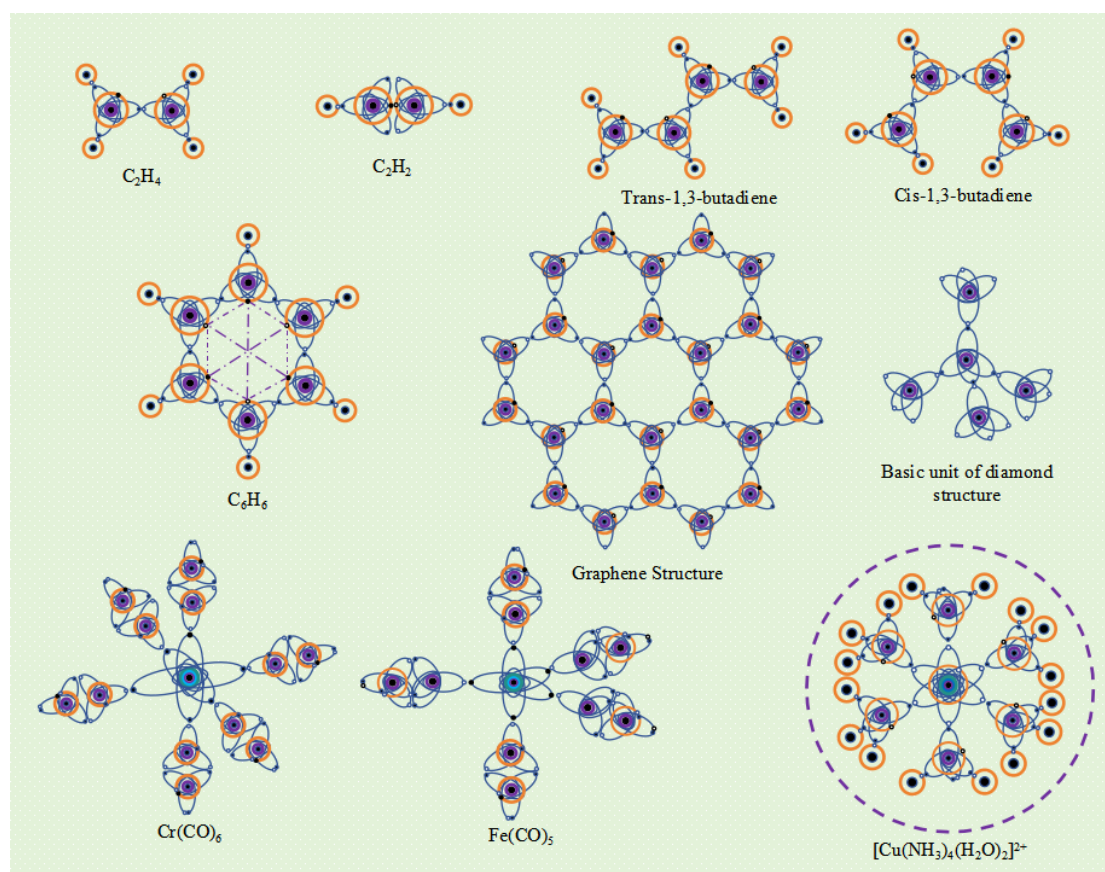


Figure 9. Structures of Several Special Covalent Molecules (Electronic Orbital Formulas).

Transition metal elements are less active than alkali and alkaline-earth metal elements, and electrons outside the atomic nucleus are less likely to become free electrons. Therefore, if a transition metal element has single electron orbits in its valence electron configuration, these single electron orbits can form covalent bonds with the single electron orbits of other atoms. For example, the outermost layer of Cr atom has 6 electrons, and the excited valence layer electron configuration is V_{6-12} , which is a regular octahedral structure. When the CO molecule is in an excited state, the valence layer electronic configuration of carbon atoms changes to V_{4-5} , and one single electron orbital of the Cr atom forms a covalent bond with one single electron orbital of the carbon atom. Each of the six single electron orbits of the Cr atom can bond with a CO molecule, forming the $Cr(CO)_6$ molecule (Figure 9).

The metal element Fe has 8 electrons in its outermost shell. In an excited state, 3 electrons become free and the resulting ion is called Fe^{3+} , leaving 5 electrons in the outermost shell, which can form a V_{5-9} electron configuration. When Fe atoms are mixed with CO molecules, 3 CO molecules receive 3 electrons from the Fe atom to become 3 CO^- ions. These CO^- ions then combine with Fe^{3+} ions under

electrostatic attraction, releasing energy that further excites the O and C atoms within the CO⁻ ions, changing their valence electron configurations to V₇₋₆ and V₄₋₆, respectively, and forming three β-bonds between the O and C atoms. Each C atom also has an elliptical single-electron orbital that forms 3 α-bonds with the three coplanar elliptical single-electron orbits in the V₅₋₉ electron configuration of the Fe atom. For CO molecules that have not acquired electrons, the bonding between the C and O atoms remains unchanged, but in the excited state, the valence electron configuration of the C atom changes to V₄₋₅, gaining an additional elliptical single-electron orbital. Two such excited CO molecules form two α-bonds with the two single-electron orbits in the V₅₋₉ electron configuration of the Fe atom that are perpendicular to the planar triangle. In this way, five CO molecules combine with one Fe atom to form a Fe(CO)₅ molecule (Figure 9). The Fe(CO)₅ molecule contains both ionic bonds between the Fe³⁺ ion and the three CO⁻ ions, and five covalent bonds between the Fe atom and C atom, making it a special compound that possesses both ionic and covalent bonds.

The electron arrangements of Group VIII elements, Group IB elements, lanthanide and actinide elements have special characteristics, and the arrangement of their inner and outermost electrons usually needs to be adjusted according to the energy state of the atomic system. When the number of outermost electrons changes, such as losing n electrons, the configuration of the valence layer electrons needs to be adjusted compared to the atoms with the previous n atomic numbers. For example, when the Cu atom of element 29 loses two electrons to become Cu²⁺, its number of electrons outside the nucleus is the same as that of element 27 Co, and the valence layer electrons are both 7. In the excited state, Cu²⁺ can form the V₇₋₁₃ electronic configuration, while the N atom in NH₃ and the O atom in H₂O form the V₅₋₈ and V₆₋₇ electronic configurations, respectively. Four excited NH₃ molecules (with N atom having an ellipsoidal single electron orbital) and two excited water molecules (with O atom having an ellipsoidal single electron orbital) form six covalent bonds with excited Cu²⁺ (with six ellipsoidal single electron orbits), forming stable [Cu (NH₃)₄ (H₂O)₂]²⁺ complex ions (Figure 9).

Some non-metallic elements also undergo changes in their valence layer electronic configuration after obtaining electrons. For example, when excited, the B atom forms a V₃₋₄ electronic configuration and can form a BH₃ covalent molecule with three hydrogen atoms. But under certain conditions, after B atom captures an electron from one of the hydrogen atoms, its valence layer electron configuration changes to V₄₋₆, forming two ellipsoidal single electron orbits. BH₃ becomes (BH₂)⁻ ion, and one hydrogen atom becomes a hydrogen ion (H⁺). The (BH₂)⁻ ion as a whole is negatively charged, and its ellipsoidal single electron orbit exhibits a negative charge property. Therefore, two (BH₂)⁻ ions can tightly bind with the electrostatic attraction between two hydrogen ions H⁺ to form diborane molecules (B₂H₆) (Figure 10a). This type of compound is called a hydrogen bridge compound, which combines two negative ions with single electron orbits through electrostatic attraction using hydrogen ions (protons).

3.2.3. Polarity of Molecules and Intermolecular Forces

The atoms that make up a molecule form different spatial structures due to their different valence electron configurations. If the electronic configuration space structure of the entire molecule is uniformly symmetrical, then the charge distribution of the entire molecular system is uniform, and the molecule exhibits non-polar characteristics, which we call non-polar molecules. If the electronic configuration space structure of a molecule is asymmetric, non-uniform, or symmetric but non-uniform, then the charge distribution of the entire molecular system is non-uniform, and the molecule exhibits polar characteristics, which we call polar molecules. In polar molecules, they exhibit negative charge properties in directions with more charge distribution and positive charge properties in directions with less charge distribution.

Taking hydrogen chloride (HCl) molecules as an example, chlorine atoms have 7 valence electrons, while hydrogen atoms only have 1 valence electron. The distribution of "electron orbital dynamic entities" in the molecular system is significantly uneven, and the charge distribution is significantly more in the direction of chlorine atoms than in the direction of hydrogen atoms.

Therefore, HCl molecules are polar molecules. The chlorine molecule (Cl_2) has two identical chlorine atoms, and due to the completely uniform and symmetrical spatial distribution of the valence layer "electron orbital dynamic entity" of the two chlorine atoms, the charge distribution is uniform and exhibits non polarity (Figure 10c).

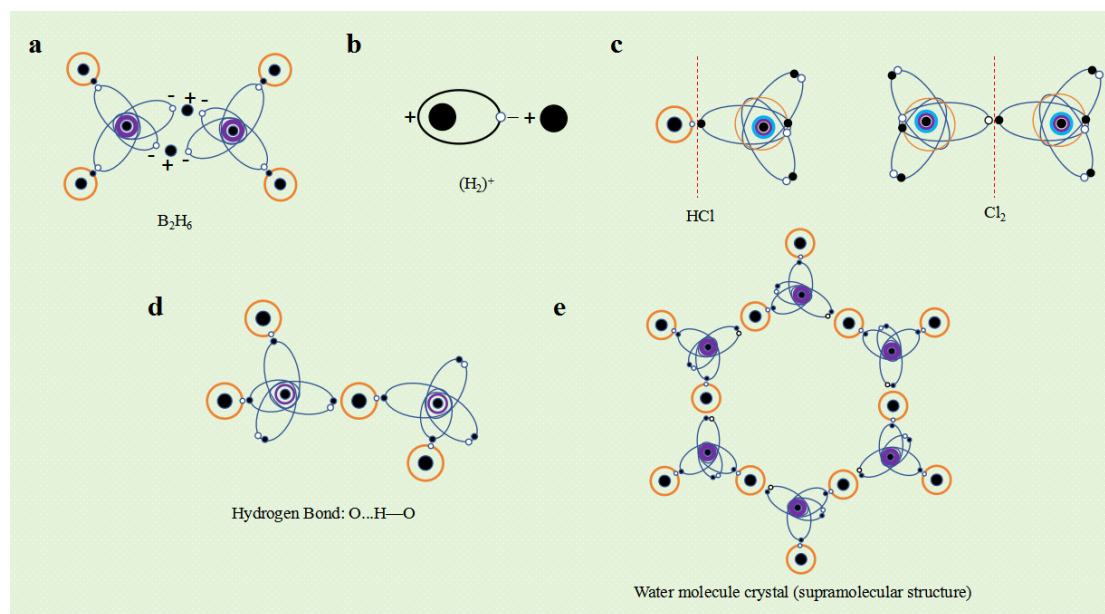


Figure 10. Atomic polarity, molecular polarity, hydrogen bond and supramolecular structure A: Hydrogen bridged compound (B_2H_6); B: Hydrogen molecular ion (H_2^+); C: Polar molecule HCl and non-polar molecule Cl_2 ; D: The hydrogen bonding structure between water molecules; E: Supramolecular Structure of Water Molecular Crystals.

The simplest atom is a hydrogen atom, where an electron orbits a proton in a circular or elliptical motion. Observing over a period of time, a hydrogen atom is a dynamic entity with a spherical or ellipsoidal electron orbit surrounding a proton. However, in the case of instantaneous observation, hydrogen atoms can be regarded as an electric dipole composed of protons and electrons at any moment. In the direction close to the electron, hydrogen atoms have a negative charge property, while in the direction close to the proton, hydrogen atoms have a positive charge property. Therefore, hydrogen atoms exhibit polarity under instantaneous observation. The hydrogen molecular ion (H_2^+) is actually the product of the instantaneous electrostatic attraction between hydrogen atoms and hydrogen ions (H^+) in a polar state (Figure 10b).

Polarized molecules (HX) containing hydrogen atoms are usually positively charged in the direction of the hydrogen atom, and correspondingly negatively charged in the direction of other atoms. The electrostatic attraction between polar molecules ($\text{HX} \cdots \text{HX}$) occurs between the hydrogen atom and other atoms, which is called hydrogen bonding. Since hydrogen atoms have only one electron outside the nucleus, the molecular charge distribution formed by their combination with other atoms is the most uneven, resulting in a higher polarity. Hydrogen bonds have stronger electrostatic attraction than other polar molecules. Figure 10d illustrates the hydrogen bonds between water molecules.

Since the overall effect of dynamic entities is not only related to the periodic motion speed but also to the observation distance, the overall effect of electronic orbital dynamic entities becomes more pronounced when the distance between molecules is far, and less pronounced when the distance between molecules is close. When the distance between molecules is very close, the instantaneous relationship between electrons and atomic nuclei exhibits an electric dipole mode, and the instantaneous interaction between molecules is manifested as the interaction between dipole moments. Due to the inherent polarity of polar molecules, the dynamic electronic orbits of non-polar

molecules are attracted to the positive charge side of polar molecules and tilt, resulting in induced dipole moments. The so-called dispersion force, induction force, and orientation force are all short-range forces, manifested as attractive forces between molecules at close distances, with a range of only a few picometers. These intermolecular forces are collectively referred to as van der Waals forces, which are essentially electrostatic forces generated by molecular or atomic polarity.

Just like metal elements and ionic compounds can form crystal structures, there is electrostatic attraction between molecules, which can also form molecular crystal structures. Due to the spatial structure of molecules and the specific valence electron configurations of the atoms that make up the molecules, the electrostatic attraction between molecules exhibits significant directionality and saturation. The polarity or instantaneous polarity between molecules generates electrostatic attraction, causing molecules to condense and form different molecular crystals or supramolecular structures. For example, water molecules interact with each other through hydrogen bonds to form water molecule crystals (Figure 10e).

When the energy state of the molecular system is low, the volume of the "dynamic entity of electron orbits" outside the ground state atomic nucleus is small, manifested as the small volume of atoms and molecules, which can tightly bind and condense into a solid "molecular cluster". When the energy state is high, the volume of atoms and molecules is relatively large, and the binding between atoms and molecules is not very tight, resulting in a condensed "molecular cluster" that appears as liquid. When the energy state is higher, the volume of atoms and molecules becomes larger, the bonding between atoms and molecules becomes less tight, the cohesion between molecules becomes weak, and the molecular aggregates appear in a gaseous state. The change of a substance composed of molecules from solid to liquid and gas is a manifestation of the energy state change of the molecular system, which is usually referred to as physical change. The change in state of matter usually refers to the change in the composition of substances composed of the same molecular aggregates. If molecules of different substances are mixed together, chemical changes may occur simultaneously with the change in the state of matter.

3.2.4. The Physical Mechanism of Chemical Reactions

The essence of a chemical reaction is the formation of new ionic or covalent bonds between different atoms that make up a substance molecule, the formation of new substance molecules from different atoms, or the formation of different molecular structures from molecules composed of the same atoms. At normal temperature and pressure, the chemical reaction in which different types of substance molecules mix to form new substance molecules is called spontaneous chemical reaction. A chemical reaction that cannot occur spontaneously at room temperature and pressure and requires conditions such as heating, pressure, or light exposure is called a non spontaneous chemical reaction.

Mixing different types of liquid molecules, or mixing between liquids and solids, or between liquids and gases, is prone to spontaneous chemical reactions. Usually, acid-base reactions, ion exchange reactions, and so-called coordination reactions are spontaneous chemical reactions. In acid-base reactions, such as mixing hydrochloric acid and sodium hydroxide solutions, hydrogen ions (H^+), chloride ions, sodium ions, and hydroxide ions (OH^-) mix together, and positive and negative ions recombine under the interaction of electrostatic attraction. H^+ and OH^- attract each other under electrostatic attraction. The excess electron outside the oxygen nucleus in the OH^- is captured by H^+ , and the valence layer electron of the oxygen atom is lost, forming an elliptical shape and becoming a single electron orbit. The H^+ obtains an electron and becomes a hydrogen atom with a single electron orbit. The single electron orbits outside the hydrogen nucleus can form covalent bonds with the single electron orbits of oxygen atoms in OH^- , generating H_2O molecules.

Sodium chloride solution is a homogeneous mixture of sodium ions and chloride ions (Cl^-), while silver nitrate solution is a homogeneous mixture of nitrate ions and silver ions (Ag^+). When sodium chloride solution and silver nitrate solution are mixed, these four ions will also be evenly distributed, which means that recombination will occur between these ions. Cl^- combines with Ag^+ under electrostatic attraction to form $AgCl$, which precipitates due to its insolubility in water.

Sodium carbonate dissolves in water and dissociates into carbonate ions (CO_3^{2-}) and sodium ions. If hydrochloric acid is added to a sodium carbonate solution, a large amount of hydrogen ions (H^+) are dissociated from hydrochloric acid in water. H^+ combines with CO_3^{2-} under electrostatic attraction, and H^+ obtains an electron from CO_3^{2-} to form a hydrogen atom. Two hydrogen atom single electron orbits form two covalent bonds with two ellipsoidal single electron orbits of oxygen atoms, generating H_2CO_3 . At room temperature and pressure, the H_2CO_3 structure is unstable and spontaneously forms CO_2 and H_2O .

Copper sulfate solution contains Cu^{2+} and SO_4^{2-} . When excessive ammonia water is added, due to the strong polarity of NH_3 molecules, H_2O molecules can dissociate into H^+ and OH^- in aqueous solution. Cu^{2+} and OH^- combine through electrostatic attraction to form $\text{Cu}(\text{OH})_2$ precipitate. The released energy will excite some atoms, transforming double electron orbits into single electron orbits and changing the valence layer electron configuration. For example, the valence layer electron configuration of Cu^{2+} is excited to V_{7-13} electron configuration, the valence layer electron configuration of N atoms in NH_3 is excited to V_{5-8} , and the valence layer electron configuration of O atoms in water molecules is excited to V_{6-7} . Therefore, Cu^{2+} can form 6 covalent bonds with 4 NH_3 and 2 H_2O , generating stable $[\text{Cu}(\text{NH}_3)_4(\text{H}_2\text{O})_2]^{2+}$ ions that combine with SO_4^{2-} to form $[\text{Cu}(\text{NH}_3)_4(\text{H}_2\text{O})_2]\text{SO}_4$.

These spontaneous chemical reactions have one thing in common, which is that after the reactants are mixed, there is electrostatic interaction between ions, and different ions recombine to form a new molecular system. The potential energy decreases, releasing energy and forming more stable ionic compounds, or exciting the valence layer electronic configuration change of atoms in ions or polar molecules (from double electron orbits to single electron orbits), and then forming stable covalent compounds.

If different substance molecules cannot be combined by electrostatic attraction after mixing, and the atoms that make up the molecules cannot form new bonds by spin magnetic force, then spontaneous chemical reactions will not occur. Usually, additional energy needs to be provided, such as heating, pressure, or light, in order for chemical reactions to occur. These additional energies increase the potential energy of molecules and atomic systems, breaking existing chemical bonds or transforming double electron orbits into single electron orbits, changing the electronic configuration of the atomic valence layer, thus allowing new bonds to form between atoms and form new compounds.

For example, in the valence layer electronic configuration of a nitrogen atom in nitrogen gas, three single electron orbits form three covalent bonds with three single electron orbits of another nitrogen atom. The two hydrogen atoms in a hydrogen molecule also form covalent bonds with two single electron orbits (Figure 8). Due to the lack of ionic interactions and single electron orbits between covalent molecules nitrogen and hydrogen, spontaneous chemical reactions do not occur at room temperature and pressure. Under high temperature and pressure, nitrogen and hydrogen molecules enter a high-energy state where the covalent bonds within the molecules are broken. Subsequently, the single-electron orbits of nitrogen atoms can form covalent bonds with the single-electron orbits of hydrogen atoms, leading to the production of ammonia.

4. Discussion

Based on the basic physical fact that electrons move in a circular or elliptical motion around the nucleus, this paper proposes a dynamic entity model of electron orbit. In this model, the shape of the electron orbit is the dynamic entity formed by the trajectory of electron movement, and the orbits of the same electronic layer are degenerate orbits with the same orbital radius and energy level. The spatial distribution of these degenerate orbits constitutes a uniformly symmetric spatial configuration of electron orbits. Therefore, in the electronic orbital space configuration, there is no concept of electron subshells as found in quantum mechanics, nor is there the issue of energy level crossing. The current electron cloud model based on quantum mechanics has successfully predicted many phenomena mathematically. However, its physical imagery—such as the shape of electron probability distributions—deviates intuitively from the trajectories of particles under central forces

in classical mechanics, often leading to pedagogical difficulties. In contrast, the electron orbital dynamic entity model proposed in this paper starts directly from the physical picture of electrons undergoing periodic motion under the Coulombic force of the nucleus, visualizing the time-averaged effect as dynamic entities with well-defined geometric shapes. This model not only aligns more closely with classical physical intuition but also avoids the complexity of tracking instantaneous positions and momenta, allowing us to focus on the overall spatiotemporal characteristics of electron motion. This approach aims to provide a conceptually more coherent and pedagogically more accessible physical imagery for atomic structure.

Based on the fundamental fact that there is electrostatic attraction between electrons and atomic nuclei, we propose the viewpoint that electron spin originates from the motion of electrons around the nucleus: the motion of electrons around the nucleus is similar to the motion of the moon around the earth, and the rotation period of electrons is the same as that of electrons around the nucleus. On this basis, the concept of electron spin quantity and the theory of electron spin quantity field were further proposed, revealing the physical mechanism of electron spin pairing and the quantization law of the mutual rotation motion of spin paired electrons. Quantum mechanics believes that the spin of electrons is an intrinsic property of electrons, avoiding the source of electron spin and the physical mechanism of electron spin pairing, making electron spin the most mysterious so-called pure quantum phenomenon. Our research has completely uncovered the mystery of electron spin, proving that electron spin is not inexplicable and mysterious, but can be deeply understood and interpreted through the principles of classical physics. This disruptive theoretical breakthrough undoubtedly brings new thinking and research directions to the physics community, marking a big step forward in our understanding of the microscopic world.

Based on the dynamic entity model of electronic orbits and the theory of electron spin, we propose the theory of electronic orbital spatial configuration, which solves the problem of multi electron atomic systems and obtains an accurate expression for the energy of the electron layer. For multi electron atomic systems, people usually use Slater's formula to calculate the energy of the electron layer, which introduces the concept of "shielding coefficient". The so-called shielding effect refers to the shielding effect of inner layer electrons on outer layer electrons, for example, when the inner layer electrons are close to the atomic nucleus, they will cancel out the positive charge of the atomic nucleus. However, Slater's "shielding coefficient" not only includes the shielding effect of inner layer electrons, but also the interaction between electrons in the same electron layer. Obviously, this coefficient does not fully conform to the concept of shielding effect. We have revised Slater's concept of "shielding coefficient", which only includes the shielding effect of inner electrons on outer electrons, and no longer includes the interaction between electrons in the same electron layer. We divide the interaction between electrons in the same electron layer into two parts: one is the mutual repulsion between dynamic entities in the electron orbit, and the other is the mutual attraction between spin paired electrons. The interaction between these two parts of electrons in the same electronic layer precisely forms a specific electronic orbital spatial configuration. Therefore, the energy of the electron layer can be expressed as the sum of three parts: "electron orbital energy level", "electron orbital spatial configuration potential energy", and "electron spin pairing energy". Based on the experimental data of atomic ionization energy, the "shielding coefficient" of different electron layers, the potential energy of electron orbital spatial configuration, and the electron pairing energy can be estimated. Our theory is very intuitive for handling multi electron atomic systems, with extremely simple methods and minimal computational complexity. This indicates that we have thoroughly solved the long-standing problem of multi electron atomic systems in the field of quantum chemistry.

Based on the principle of minimum energy and the theory of electronic orbital spatial configuration, we rearranged the extranuclear electrons of all elements in the periodic table. In quantum mechanics, the arrangement of extranuclear electrons around atomic nuclei is highly complex, with frequent exceptions that do not conform to established rules, and there are also errors in the arrangement of inner-shell electrons where the number is odd. In quantum mechanics, the

electron configurations of many elements, especially those in Group VIII, Group 1B, and the lanthanides and actinides, are complex, often exhibiting numerous exceptions that do not conform to the established rules of arrangement. Additionally, there are errors where the inner electron configurations have odd numbers. We have abandoned the concept of electron subshells in quantum mechanics, corrected the erroneous electron configurations, and established a new method for electron arrangement. In this method, the extranuclear electron configuration of each element follows the same set of rules without any exceptions. Our method for determining the extranuclear electron configuration of elements is characterized by its simplicity and accuracy, which is crucial for correctly understanding the chemical properties of elements.

Based on the theory of electron spin pairing and spatial configuration of electron orbits, we have solved the problem of multi atomic molecules, revealed the nature and formation mechanism of chemical bonds, and explained and predicted the spatial structure of covalent compounds. The existing VB and MO theories are powerful computational tools. However, in terms of physical imagery, there exists a conceptual tension in VB theory between the notion of "orbital overlap" and the inevitably increased Coulombic repulsion between electrons, which requires further clarification. Meanwhile, the MO theory's picture of "electrons delocalized over the entire molecule" is not intuitively consistent with the fundamental physical reality that electrons are strongly bound to individual atomic nuclei. The theory proposed in this paper attempts to return to a more foundational physical picture: chemical bonds arise from magnetic interactions between unpaired electron orbital dynamic entities (covalent bonds) or purely electrostatic interactions (ionic bonds, metallic bonds). Within this framework, the formation, directionality, and strength of bonds are directly determined by the spatial configuration (V_{n-m}) of electron orbits and the mode of spin pairing (α/β bonds), thereby offering an alternative explanatory scheme for the nature of chemical bonds based on classical electromagnetic interactions and geometric arrangement. For instance, in the case of the benzene molecule, this theory can naturally and logically derive its highly stable hexagonal ring structure without invoking the special hypothesis of a "delocalized large π bond". Instead, it relies solely on specific forms of orbital magnetic pairing between carbon atoms (Type I and Type II α bonds), providing an alternative explanation for conjugated systems based on localized interactions with a clearer physical imagery.

Our theory can explain the formation and structure of coordination compounds (complexes) of transition elements. The VB theory holds that complexes are formed by the central atom or ion providing empty orbitals, and the ligand molecule or ion providing lone electron pairs, which are bound through coordination bonds. However, the concept of empty orbitals based on quantum mechanics theory has serious problems. The so-called "empty orbitals" contain no electrons, which practically means there are no orbits at all, rendering such empty orbitals entirely fictitious. Additionally, according to the theory of quantum mechanics, each atom has an infinite number of empty orbitals, implying that it is not solely the central atoms or ions of transition elements that can provide empty orbitals. The coordination field theory and crystal field theory suggest that the lone pair electrons provided by the ligand will affect the energy distribution of the central atom's d orbital, causing the d orbital to split into different energy levels. However, as previously pointed out, the concept of electron subshells in quantum mechanics is flawed, and the so-called d -orbitals do not exist. Furthermore, the process of so-called d -orbital energy level splitting and coordination bond formation involves changes in the shape and distribution of electron clouds, as well as the overlap of electron clouds, which is very complex and far fetched. According to our theory, when the central atoms or ions of transition elements and the central atoms of polar molecules are excited, the electronic configuration of the valence layer changes, and the double electron orbits become single electron orbits, forming covalent compounds (such as the $\text{Cr}(\text{CO})_6$). Alternatively, in the excited state, free electrons in transition metals are captured by polar molecules, causing changes in the valence electron configuration of the central atom or ion, converting double-electron orbits into single-electron orbits, and then forming covalent compounds with ionic bond characteristics (such as $\text{Fe}(\text{CO})_5$). The metal ions of transition elements are prone to form complexes with polar molecules

because after losing electrons, the valence layer electronic configuration of these transition elements will be adjusted according to the energy state of the atomic system. At the same time, the release of energy during the reaction process or under additional energy conditions will excite the central ions of transition elements and the central atoms of polar molecules, causing a change in their valence layer electronic configuration, and the double electron orbits will become single electron orbits, thereby forming covalent bonds. We can determine the molecular structure of the complex based on the valence electron configuration of the central ion of the transition element in the excited state. We can explain and predict the formation and structure of complex coordination compounds solely based on the changes in the valence electron configuration of the central atom or ion of the transition element, as well as the central atom of the ligand. This simplicity and accuracy suggest the correctness of our theory.

VB theory and MO theory are difficult to explain the formation and structure of hydrogen bridged compounds. Currently, the most satisfactory explanation for hydrogen-bridged compounds is considered to be the "three-center two-electron bond" model¹⁵. This model suggests that the B atom in diborane (B_2H_6) adopts unequal sp^3 orbitals, one of which is an empty orbital. However, the so-called empty orbit does not exist, and using the concept of empty orbit to explain the formation of chemical bonds is very far fetched. According to our theory, the key to the formation of hydrogen bridged compounds is that atoms with three valence electrons, such as B atom, can capture electrons from the hydrogen atom when reacting with it, causing the valence electron configuration to change from V_{3-4} to V_{4-6} and the hydrogen atom to become a hydrogen ion. Excited B ions form negatively charged BH_2 atomic clusters with hydrogen atoms, where two ellipsoidal single electron orbits combine with another negatively charged BH_2 atomic cluster through electrostatic attraction to form B_2H_6 . Our theory reveals that the essence of hydrogen bridge bonds is to connect two negatively charged atomic clusters through hydrogen ions, which is a new breakthrough in the understanding of hydrogen bridge compounds.

VB theory and MO theory cannot explain the changes in solid, liquid, and gas states of material molecules, and cannot directly explain the crystal and supramolecular structures of molecules. Additionally, these theories are also difficult to explain chemical reactions. Our theory can not only explain the physical principles of molecular polarity and atomic instantaneous polarity to explain the changes in state of matter and the structure of molecular nanocrystals and supramolecular structures, but also explain chemical reactions and the formation of molecular structures from the relationship between atomic energy states and valence layer electron spatial configurations. When we explain the physical changes of matter and the chemical changes of molecules, the theoretical framework we rely on is unified and the logical framework is also consistent, which strongly proves the high inherent unity and universality of our theory.

5. Conclusion

This paper establishes a comprehensive theoretical system of atomic and molecular structure based on the fundamental principles of classical physics. This system, supported by the Dynamic Entity Model, the Electron Spin Theory derived from orbital motion, and the Spatial Configuration Theory of Electron Orbits, achieves for the first time a logically unified explanation within a classical framework—spanning from atomic structure to molecular properties (structure, chemical bonds, and reactivity).

The core advantages of this theory are as follows:

1. Conceptual Intuitiveness and Self-consistency: All concepts are rooted in classical mechanics and electromagnetism, avoiding the introduction of notions such as "probability waves" and "intrinsic spin", which are detached from classical intuition. The resulting physical picture is clear, providing a powerful tool for chemical education and interdisciplinary understanding.

2. Predictive Power and Testability: The theory deduces a series of unique and strongly falsifiable predictions, the most prominent being the novel electron arrangement rule that **the inner-**

shell electron count of all elements must be even (Table 1). This offers a clear target for experimental verification via spectroscopy and other methods.

3. Computational Feasibility and Application Potential: As demonstrated in Section 2.5 using the boron atom as an example, this theory provides a systematic computational framework for decomposing macroscopically observable data (ionization energies, spectra) into microscopic components with clear physical meanings (shielding energy, configuration energy, pairing energy). This feature indicates significant potential application value in the following areas:

Intuitive Elucidation of Chemical Reaction Mechanisms: Based on changes in electron orbital configurations and transformations between bond types (α/β bonds), this theory holds promise for offering more concrete physical imagery and predictive models for the microscopic pathways of chemical reactions (e.g., concerted reactions, catalytic cycles) than current theories.

Rational Design of Molecular and Material Structures: The theory directly links the energy state of atoms (V_{n-m} configuration) to the final spatial geometry of molecules. This makes it possible, starting from the atomic level, to **predict or even design molecular configurations with specific functions** (e.g., active sites of catalysts, chirality of drug molecules) and material crystal structures. It provides a novel, classically-based, visual simulation approach for molecular modeling and materials science.

Property Prediction: By calculating the total configuration potential energy and bond energy distribution of molecular systems, this theoretical framework is expected to develop new methods for predicting molecular stability, spectral properties, and reactivity.

While the current theoretical system based on quantum mechanics is widely applied, it still faces challenges regarding its conceptual foundations and unification. The theory developed in this paper is a complete "**alternative theory**" based on the classical physics framework. It demonstrates that the logical system of classical physics retains powerful, yet underexplored, explanatory and predictive potential at the microscopic scale. We anticipate that this theory will not only stimulate new reflections in fundamental physics but also serve as an **intuitive and powerful tool** to open up new research paradigms in applied fields such as chemical synthesis, drug design, and materials exploration, while subjecting itself to the most rigorous experimental and computational scrutiny.

Data Availability Statement: No Data associated in the manuscript.

Acknowledgments: We sincerely thank Professor Guihua Lu for pointing out the errors and omissions in the preprint of the manuscript, which helped us make revisions.

References

1. Rutherford, E. The scattering of alpha and beta particles by matter and the structure of the atom. *Philosophical Magazine* **21**, 669-688 (1911).
2. Bohr, N. On the Constitution of Atoms and Molecules. *The London, Edinburgh, and Dublin Philosophical Magazine and Journal of Science* **26**, 1-25 (1913).
3. Sommerfeld, A. Zur quantentheorie der spektrallinien. *Annalen Der Physik* **5**(18), 5-9(1916).
4. Mermin, D. N. What's Wrong with this Pillow? *Physics Today* **42**(4), 9-11(1989).
5. Zeng, J.Q., Zeng, T. H. Study on the diffraction-like and interference-like mechanisms of particle flow. *Applied Physics Research* **5**(2), 157-172(2023).
6. Lewis, G.N. The Atom and the Molecule. *J. Am. Chem. Soc.* **38**, 762-785 (1916).
7. Heitler, W., London, F. Wechselwirkung neutraler Atome un homöopolare Bindung nach der Quantenmechanik. *Z. Phys.* **44**, 455-472(1927).
8. Pauling, L. The Shared-Electron Chemical Bond. *Proc. Natl. Acad. Sci. USA* **14**, 359-362(1928).
9. Slater, J.C. Molecular Energy Levels and Valence Bonds. *Phys. Rev.* **38**, 1109-1144(1931).
10. Mulliken, R.S. The Assignment of Quantum Numbers for Electrons in Molecules. I. *Phys. Rev.* **32**, 186-222(1928).
11. Hund, F. Zur Frage der Chemischen Bindung. *Z. Phys.* **73**, 1-30(1931).

12. Bethe, H. A., & Bright, A. W. The theory of the crystal structure of transition metal salts. I. Magnetic and spectral properties. *Proceedings of the Royal Society of London. Series A, Mathematical and Physical Sciences* **126**(802), 238-265(1930).
13. Jørgensen, C. K. On the constitution of complex salts. I. Electronic theory. *Acta Chemica Scandinavica* **6**, 825-836(1932).
14. Pauling, L. The nature of the chemical bond—application of results obtained from the quantum mechanics and from a theory of paramagnetic susceptibility to the structure of molecules. *Journal of the American Chemical Society* **61**(6), 1466-1475(1939).
15. Zeng, J.Q. Classical physical mechanism of quantum production and its explanation for hydrogen atom structure and photoelectric effect. *Physics Essays* **34**(4), 529-537(2021).
16. Zeng, J.Q. Classical physics derivation of quantization of electron elliptical orbit in hydrogenlike atom. *Physics Essays* **35**, 147-151(2022).
17. Zeng, J.Q., Zeng, T.H. The Great Tao Model: The Theory of Fundamental Particles and Their Interactions. *in submission*.
18. Xu, G.X., Wang, X.Y. Material structure. *Second edition. Science Press* (2010)

Disclaimer/Publisher's Note: The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.