

UNIT 4 : CHEMICAL BONDING

Chemical Bonding: The Force of Attraction in the Molecular World

Introduction

At the heart of the vast diversity of substances and the myriad of chemical reactions lies a fundamental concept: chemical bonding. This is the force that holds atoms together allowing them to form molecules and compounds. Chemical bonds are crucial for the existence of everything, from the air we breathe to the DNA in our cells. Let's delve into the types and nature of chemical bonds.

1. What is Chemical Bonding?

Chemical bonding refers to the force of attraction between atoms or ions. This force arises due to the need for atoms to achieve a stable electron configuration, often resembling that of noble gases. The type and strength of these bonds determine the chemical and physical properties of substances.

2. Types of Chemical Bonds:

- Covalent Bonds:
 - Formation: When two atoms share one or more pairs of electrons.
 - **Characteristics:** Found in non-metal compounds, can be polar (unequal sharing of electrons) or non-polar (equal sharing of electrons).
 - Example: Water (HzO), where oxygen shares electrons with two hydrogen atoms.
- Ionic Bonds:
 - Formation: Result from the transfer of electrons from one atom (which becomes a cation) to another (which becomes an anion). The electrostatic attraction between the oppositely charged ions forms the bond.
 - **Characteristics:** Usually formed between metals and non-metals, resulting in crystalline solids at room temperature.
 - **Example**: Table salt (NaCl), where sodium donates an electron to chlorine, forming Na* and Cl- ions.
- Metallic Bonds:
 - Formation: Occur between metal atoms.
 - **Characteristics:** Metal atoms release their outer electrons, which are then free to move throughout the entire metal lattice, forming a 'sea' of delocalized electrons.
 - **Example**: Copper (Cu) and its ability to conduct electricity due to the mobility of its delocalized electrons.

3. Other Types of Bonds and Interact1ons:

• **Hydrogen Bonds:** A type of dipole-dipole interaction where a hydrogen atom, already covalently bonded to an electronegative atom (like oxygen or nitrogen), forms an electrostatic attraction with another electronegat1ve atom.

• Van der Waals Forces: Weak forces arising from instantaneous dipoles in molecules. IncludesLondon dispersion forces and dipole-dipole interactions.

4. Bond Strength and Length:

The strength of a bond is related to the bond energy, which is the energy required to break a bond. Generally:

- Triple bonds > Double bonds > Single bonds (in terms of strength and energy).
- Single bonds > Double bonds > Triple bonds (in terms of length).

Conclusion:

Chemical bonding underlies the structure and reactivity of all substances around us. Whether forming the crystal lattice in gemstones or ensuring the right fit between enzymes and substrates in our body, chemical bonds play an indispensable role in the universe's molecular dance. Understanding them is foundation t fields ranging from medicinal chemistry to materials science.

1.1 Indian Chemistry: - Philosophy of atom by Acharya Kanad. Indian Chemistry: Acharya Kanad and the Philosophy of the Atom

Introduction

Long before the modern atomic theory emerged in the West, ancient India had already delved deep into the concept of the atom. One of the foremost proponents of this early atomic theory was Acharya Kanad, an ancient Indian philosopher and sage. His profound ideas paved the way for a more detailed understanding of the microscopic world in the Indian subcontinent.

1. Acharya Kanad: An Overview

Acharya Kanad, also known as Kashyap, lived around the 6th century BCE. He was a philosopher and the founder of the Vaisheshika school of philosophy, one of the six classical schools in Hindu philosophy. The name "Kanad" is believed to have originated from "Kana," which means "particle" or "atom" in Sanskrit, underscoring his deep association with atomic theory.

2. The Philosophy of the Atom

- Anu (Atoms): According to Acharya Kanad, the universe is composed of minute particles called "anu" or atoms. These atoms are indivisible, indestructible, and eternal. They are the fundamental building blocks of matter.
- **Combination and Division:** Kanad posited that these atoms could combine in various ways to form complex structures, called "dwinuka" (binary molecules). When the right conditions prevailed, these combinations could be reversed, breaking down complex structures into their atomic constituents.

• **Properties:** Each atom possessed its own set of properties. However, when atoms combined, new properties emerged that were different from those of the individual atoms. This notion aligns with the modern understanding that properties of molecules can be distinct from those of their constituent atoms.

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3. Contribution to Indian Thought

Acharya Kanad's atomic theory was groundbreaking and formed the foundation for the Vaisheshika school's more extensive framework. This philosophy delved deep into the nature of matter classifying entities into various categories, establishing causality, and providing insights into the universe's mechanics.

While the Vaisheshika atomic theory and modern atomic theory have distinctions, Kanad's contributions represent a significant leap in abstract thinking and the quest to understand the universe's fundamental nature.

4. Legacy and Influence

The Vaisheshika school, along with Kanad's atomic philosophy, influenced other schools of Indian thought and played a critical role in shaping early Indian scientific and philosophical discourse. The ideas resonated across generations, finding echoes in later texts, commentaries, and discussions.

Conclusion

Acharya Kanad's philosophy of the atom serves as a testament to the depth and breadth of ancient Indian scientific and philosophical thought. While separated by millennia, Kanad's insights and the contemporary understanding of the atom bridge the vast expanse of time, underscoring the timeless quest for knowledge and the desire to understand the universe's intricate fabric.

1.2 Electronic theory of valency: Assumptions, Chemical bonds: Types and characteristics of electrovalent bond, covalent bond, coordinate bond, hydrogen bond, metallic bond and Intermolecular forces of attraction.

Electronic Theory of Valency and Chemical Bonds: An Exploration

Introduction

The electronic theory of valency offers a foundational framework for understanding chemical bonding, explaining why atoms come together to form molecules. This theory, combined with the knowledge of atomic structure, paves the way for a deeper comprehension of various bond types and their characteristics.

1. Electronic Theory of Valency:

- Assumptions:
 - Atoms combine to achieve a stable noble gas configuration, often an octet in the outermost shell.
 - The number of valence electrons (in the outermost shell) dictates an atom's combining capacity or valency.
 - Atoms can achieve stability either by losing, gaining, or sharing electrons.

2. Chemical Bonds: Types and Characteristics:

- Electrovalent (Ionic) Bond:
 - **Formation:** Resulting from the transfer of electrons from one atom to another.
 - **Characteristics:** Typically formed between metals and non-metals. High melting and boiling points, conduct electricity in molten or aqueous state.
 - **Example**: NaCl, where sodium donates an electron to chlorine.
 - Covalent Bond:
 - **Formation**: Originates from the sharing of electrons between two non-metal atoms.
 - Characteristics: Can be polar (unequal sharing) or non-polar (equal sharing). Lower melting and boiling points compared to ionic compounds. Poor conductors of electricity.
 - **Example:** H2O, where two hydrogen atoms share electrons with an oxygen atom.
 - Coordinate (Dative) Bond:
 - Formation: A type of covalent bond where one atom provides both electrons for sharing.
 - Characteristics: Common in complex ions and certain molecules.
 - **Example**: The bond between boron and one of the hydrogens in BH(— ion.
 - Hydrogen Bond:
 - Formation: A special type of dipole-dipole interaction between a hydrogen atom (bonded to a highly electronegative atom) and another electronegative atom.
 - **Characteristics:** Weaker than covalent and ionic bonds, but stronger than regular van der Waals forces. Responsible for many physical properties, such as the high boiling point of water.
 - **Example:** Bond between water molecules in liquid HzO.

Metallic Bond:

- Formation: Occurs between metal atoms.
- Characteristics: Metal atoms release their valence electrons, forming a 'sea' of delocalized electrons. High electrical and thermal conductivity, malleable and ductile.
- **Example:** The bond within a piece of copper metal.



3. Intermolecular Forces of Attraction:

- Van der Waals Forces: Weak attractions between molecules. They can be induced dipoleinteractions or temporary fluctuations in electron distribution.
- **Dipole-Dipole Interactions:** Attractions between molecules with permanent dipoles, as seen in polar molecules.
- London Dispersion Forces: Resulting from temporary dipoles induced by fluctuations in the electron cloud of non-polar molecules.

Conclusion

The electronic theory of valency and the subsequent classification of chemical bonds provide insights into the microscopic world, explaining the myriad of substances and their diverse properties. This knowledge serves as a bedrock for fields ranging from inorganic chemistry to materials science, elucidating the structure, behavior, and interactions of matter at the atomic and molecular levels.

1.3 Molecular arrangement in solid, liquid and gases. Molecular Arrangement in Solids, Liquids, and Gases

Introduction

The three primary states of matter - solids, liquids, and gases - differ in their properties and behaviors. A significant reason behind these differences lies in the molecular arrangements within each state. Let's delve into the intricacies of these arrangements and their implications.

1. Solids:

- Arrangement: In solids, particles (which can be atoms, molecules, or ions) are packed closely together in a well-ordered, definite geometric pattern. This rigid structure is known as a crystal lattice in crystalline solids.
- **Movement:** Particles in solids vibrate about fixed positions but do not have the freedom to move around or slide past each other.
- Implications: This close, fixed arrangement results in solids having a defined shape and volume. They are not easily compressible and have high densities compared to liquids and gases.

2. Liquids:

- Arrangement: In liquids, particles are still close together, but not as tightly packed as in solids. There is no fixed pattern, and the arrangement is more random.
- **Movement**: Particles in liquids have more kinetic energy than in solids. They can slide past each other allowing the liquid to flow and take the shape of its container while maintaining a relatively fixed volume.
- **Implications:** The increased space between particles in liquids, as compared to solids, results in a slightly lower density and the ability to be slightly compressed under pressure. Liquids

can flow and have a surface tension that leads to phenomena like droplet formation.

3. Gases:

- Arrangement: In gases, particles are far apart with a lot of empty space between them. The arrangement is random and dispersed.
- Movement: Gas particles possess high kinetic energy, allowing them to move freely and rapidly in all directions. They frequently collide with each other and with the walls of their container.
- Implications: Gases have no fixed shape or volume; they expand to fill their container entirely. They are easily compressible due to the large amount of empty space between particles and have lower densities compared to solids and liquids.

Conclusion

The diverse molecular arrangements in solids, liquids, and gases account for the distinctive properties exhibited by each state of matter. Whether it's the rigidity of a solid metal block, the fluidity of water, or the expansiveness of the air we breathe, the underlying particle arrangements dictate the macroscopic behaviors we observe. Understanding these arrangements is pivotal in fields ranging from material science to meteorology, providing insights into matter's fundamental nature and behavior.

1.4 Structure of solids: crystalline and amorphous solids ,Properties of metallic solid, Unit cell: simple cubic, body center cubic (BCC) , face centre cubic (FCC), hexagonal close pack crystals.

Structure of Solids: A Journey into the Microscopic World

Introduction

Solids, with their fixed shapes and volumes, exhibit a world of microscopic intricacies. From the orderly pattern of crystalline solids to the chaotic structure of amorphous solids, the realm of solid-state chemistry offers insights into the nature and behavior of materials that shape our daily lives.

1. Crystalline vs. Amorphous Solids:

- Crystalline Solids:
 - **Structure:** Have a long-range, repetitive order of particles. They display a definite geometric pattern known as a crystal lattice.
 - **Properties:** Exhibit sharp melting points, can diffract X-rays, and often have directional properties.
 - Examples: Salt (NaCl), sugar, and diamonds.



- Amorphous Solids:
 - **Structure:** Lack a regular repeating pattern, resembling the structure of liquids on a microscopic scale.
 - Properties: Do not have a sharp melting point, instead they soften over a range of temperatures. They do not diffract X-rays in the distinct pattern seen in crystalline solids.
 - Examples: Glass, rubber, and many polymers.

2. Properties of Metallic Solids:

- **Structure:** Comprise metal atoms packed closely in geometric patterns.
- **Bonding:** Characterized by a sea of delocalized electrons that move freely around the positive metal ions.
- **Properties:** High electrical and thermal conductivity, malleability, ductility, luster, and high melting and boiling points.

3. Unit Cells: The Building Blocks of Crystals:

- Simple Cubic:
 - Description: Atoms are situated at each corner of a cube.
 - **Coordination Number**: 6 (each atom is in contact with six others).
- Body-Centered Cubic (BCC):
 - **Description**: Atoms are located at every corner and a single atom is present at the cube's center.
 - Coordination Number: 8.
- Face-Centered Cubic (FCC):
 - **Description**: Atoms reside at each corner of the cube and at the center of each face.
 - Coordination Number: 12.
 - Hexagonal Close-Packed (HCP):
 - **Description:** Consists of hexagonal layers of atoms. Each atom is surrounded by 12 others.
 - Coordination Number: 12.

Conclusion:

The microscopic structure of solids, whether crystalline or amorphous, underpins their macroscopic properties, which, in turn, dictate their applications and functionalities. Understanding these structures and their inherent patterns enables scientists and engineers to tailor materials for specific applications, from designing robust metallic alloys to crafting transparent glass for optical applications. The world of solids, thus, offers a blend of beauty, order, and practicality.



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