

Rachana Gaur¹, Ruby Jindal^{1*}, Archana Tripathi^{2*}, Harleen Kaur³,
Aparna Shekhar⁴

¹School of Basic and Applied Sciences, K.R. Mangalam University, Gurugram, Haryana, 122103, India, ²Department of Physics, Deshbandhu College (University of Delhi), New Delhi, 110019, India, ³Department of Applied Sciences, Baba Banda Singh Bahadur Engineering College, Fatehgarh Sahib, Punjab, 140407, India, ⁴Department of Chemistry, Deshbandhu College (University of Delhi), New Delhi, 110019, India

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Columbite-Structured AB_2O_6 Oxides: Understanding their Structure, Synthesis, and Potential for Future Technologies

ABSTRACT

AB_2O_6 Oxides have recently attracted significant attention these days due to their exclusive properties. These compounds exhibit a variety of structural forms with diverse characteristics. Among them, columbite-type Oxides possess an orthorhombic structure, belong to $Pbcn$ space group and exhibit D_{2h}^{14} symmetry. The structure of these material consists of AO_6 and BO_6 octahedra which are arranged in zig zag manner and have a powerful impact on the characteristics of these materials. These materials play a vital role in different applications such as microwave technology, catalysis, energy storage, electronic sensors and optical materials. From a comprehensive literature survey, this review has covered the detailed study of the structure, physical properties, synthesis techniques, and the applications of these Oxides. Subsequently, we discuss the future potential of the Columbite Oxides, with an emphasis on strategies and computational modeling to enhance their properties and performance for the future technologies.

Keywords: Columbite, thermos electric, microwave technology, computational modeling

1. INTRODUCTION

1.1 The need for columbite AB_2O_6 oxides

The global demand for eco-friendly and sustainable energy alternatives continues to grow, largely driven by concerns over global warming linked to fossil fuel consumption, which poses a serious threat to our environment and long-term survival. Among the promising renewable energy technologies, thermoelectric (TE) modules are gaining attention. Thermoelectric devices can convert waste heat originated from various sources, such as power plants, industrial processes, vehicles, and electronic devices into electricity via Seebeck effect.

However present TE materials have low efficiency, which limits their broad commercial use. Also, a large number of these materials suffers from instability at high temperatures. On the other hand, oxide materials have become gradually attractive due to their abundance, cost-effectiveness and reduced environmental impact.

Besides, certain oxides exhibit spin-related properties that offer fascinating possibilities for optimizing their behavior for multiple applications. Among them, AB_2O_6 -type oxides are notably significant due to their distinct compositions and structural variations. Exploring the impact of the structure on the properties may expand the possibilities for thermoelectric applications at high temperatures [1].

More than 50% of the input energy is wasted as heat in almost half of the industrial processes [2,3]. This energy loss occurs in many systems extended from household items like kitchen kettles to large-scale manufacturing and power generation facilities. In the same way, gasoline-powered vehicles utilize only about 25% of the fuel's energy into actual motion and associated functions, while the rest is wasted as heat through exhaust gases, engine cooling systems, friction, and other parasitic effects. A sustainable energy solution involves taking this waste heat and converting it into electricity using specific devices. These devices work on thermoelectric principles, where heat is converted directly into electrical power through Seebeck effect, a phenomenon in which a voltage is induced across a material due to a temperature difference, causing an electric current to flow. Thermoelectric (TE) modules produce electricity without any mechanical component and without releasing greenhouse gases. Power generation devices are known as thermoelectric generators

*Corresponding author: Ruby Jindal

E-mail: ruby.jindal@krmangalam.edu.in(RJ)

*Corresponding author: Archana Tripathi

E-mail: archana851@gmail.com(AT)

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(TEGs) [3]. Current TE materials face challenges such as poor thermal stability, insufficiency, deadliness, which has prompted the search for alternatives that are abundant, environment-friendly, and thermally stable at higher temperatures. Many thermoelectric (TE) materials contain elements that are limited and toxic. For instance, lead (Pb) gives rise to serious health hazards, which includes brain damage, seizures, and coma. Materials that contains lead and antimony, are particularly harmful emphasizing the urgent need of environmentally friendly alternatives, such as TE oxides, for thermoelectric applications [1]. TE oxides offer greater thermal stability compared to current TE materials, making them strong candidates for high-temperature applications [3].

As 5G technology has become everyday technology, scientists are already working on the next advancement—6G. This upcoming technology aims to provide incredibly fast data speeds, potentially in terabits per second, while supporting the growing number of connected devices in the Internet of Things (IoT). To achieve these goals, 6G will work in terahertz frequency spectrum, which will require advanced electronic components [4,5]. Mie theory explains how electromagnetic waves interact with tiny particles, especially how certain resonances occur in materials like dielectric spheres and how the wave behave at high temperatures [6]. In the terahertz range, wave propagate differently—they're more likely to scatter or reflect from the surfaces, especially when there's no direct line of sight between the transmitter and receiver [7,8]. To tackle these challenges, scientists are exploring special materials called dielectric metamaterials. These don't rely on electric currents like traditional materials but instead use the movement of electric charges within the material itself. Their performance, however, depends a lot on how much energy they lose in the process, so understanding these losses is key for 6G development [9–12].

A group of materials called columbite niobates (with the formula ANb₂O₆, where A can be Zn, Co, Mn, Ni, etc.) is gaining attention because they have useful features due to their unique structure and the different properties of the metals involved [13–17]. Among them, ZnNb₂O₆ and MgNb₂O₆ stand out for their very low energy losses at microwave frequencies (with Qxf values above 80,000 GHz), making them strong candidates for next-generation electronic components. Interestingly, even though some of these materials look and behave similarly, they still show big differences in performance—for example, between ZnNb₂O₆ and NiNb₂O₆—and scientists are still figuring out why [18,19].

2. STRUCTURE

Columbite-type compounds with the general formula AB₂O₆ exhibit a well-defined crystal structure that belongs to the orthorhombic crystal system, specifically categorized under the space group Pbcn (No. 60). In this structural framework, the A and B represent two different metal cations, each coordinated by six oxygen atoms to form AO₆ and BO₆ octahedra, respectively. These octahedra are not randomly distributed; instead, they are systematically arranged in layers, forming a unique zigzag stacking sequence that follows an -A-B-B-A pattern along the crystallographic [100] direction [20]. The AO₆ and BO₆ octahedra are connected to one another via edge-sharing, i.e. by sharing two common oxygen atoms along their edges resulting in the formation of distinct one-dimensional chains, each made up of alternating A and B octahedra. These chains repeat in a particular ABBABB sequence, and are not confined but again interconnected via corner-sharing, where they share single oxygen atoms at the corners of octahedra. This connectivity between the chains resulting a strong and continuous three-dimensional structure, giving rise to a stable structural pattern identified by AO₆–BO₆–BO₆ chains [21–27]. The spatial distribution of A and B cations within the structure is majorly determined by their ionic radii and oxidation states. Generally, the A-site cation is a divalent transition metal ion such as Mn²⁺, Fe²⁺, Co²⁺, or Ni²⁺, whereas the B-site is accommodated by pentavalent metal ion like Nb⁵⁺ or Ta⁵⁺ [20,28]. This unique combination of cation sizes and charges supports the systematic filling of the octahedral voids in the oxygen lattice, contributing to the structural stability. Structurally, the columbite-type framework is derived from the α-PbO₂ structure, where oxygen atoms are arranged in a slightly distorted hexagonal close-packed (HCP) configuration. In this derivative structure, only half of the octahedral voids created by the oxygen packing are occupied by metal cations. This partial filling gives rise to two crystallographically distinct octahedral sites, referred to as the 4c and 8d positions [29,30], which are key in determining the cation coordination and overall symmetry.

In summary, the AB₂O₆ columbite structure is characterized by a well-organized three-dimensional network composed of zigzag chains of edge-sharing AO₆ and BO₆ octahedra, interconnected through corner-sharing. This combination of geometric ordering and cation size-dependent occupancy leads to a structurally versatile and chemically robust material class, making columbite-type oxides particularly significant in solid-state chemistry and materials science [31–33].

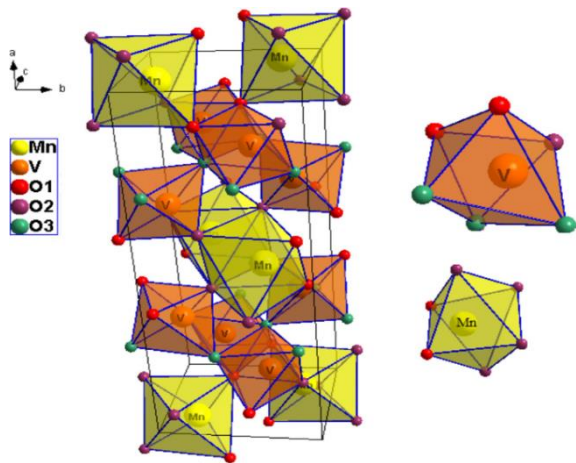


Figure 1. Structure of Columbite Oxides using DIAMOND

3. METHODS OF SYNTHESIS OF COLUMBITE-TYPE AB₂O₆ OXIDES

From previous research, various synthesis techniques have been employed to produce columbite-type AB₂O₆ oxides. These methods include conventional solid-state reactions [34,35], hydrothermal synthesis [36,37], pulsed laser deposition [38], the molten salt method [39], polymeric precursor methods [40], ceramic flux techniques [41,42], precipitation [43], and molten salt synthesis [44,45]. A selection of these methods is discussed in detail below:

High-Pressure Synthesis:

Polycrystalline MV₂O₆ (M = Mn, Co, Ni, Zn) samples were synthesized under phase-pure precursors under pressures ranging from 5.5 to 6.7 GPa at temperatures of 700–900 °C using Belt and Conac presses, with a holding time of 60 minutes followed by rapid cooling. Room-temperature X-ray diffraction (XRD) was conducted using Cu-Kα_{1,2} radiation, and structures were refined using the Rietveld method with FULLPROF, while VESTA was used for visualization [46].

Hydrothermal Method:

MnNb₂O₆ and Ca-substituted derivatives were synthesized hydrothermally by reacting NH₄(NbO(C₂O₄)₂)·(H₂O)₂·xH₂O with MnSO₄·H₂O in a 2:1 molar ratio, followed by alkali addition (LiOH, NaOH, or KOH), stirring, and heating at 240 °C for 18 hours. Products were treated with 0.5 M acetic acid to eliminate MnCO₃ impurities, then sintered at 1000 °C [47].

Sol-Gel and Solid-State Methods:

ZnNb₂O₆, NiNb₂O₆, and CoNb₂O₆ were synthesized via both sol-gel and solid-state methods. ZnO and Nb₂O₅ were calcined at 1000 °C, and NiO nanoparticles were derived

from nickel nitrate hexahydrate and ethylene glycol before calcination at 800 °C. The final columbite phases were obtained after solid-state reactions [48].

Polymeric Precursor Route:

MgNb₂O₆ was synthesized using Nb₂O₅, NaOH, and glacial acetic acid to form hydrated niobic acid, followed by complexation with citric acid. A viscous liquid formed was evaporated and annealed at varying temperatures to obtain the crystalline ceramic phase [49].

Ammonia-Assisted Synthesis:

ZnNb₂O₆ was also synthesized using ZnCl₂ in strongly alkaline conditions (pH 14) with ammonia as a mineralizer. Alternatively, ZnNb₂O₆ powders were prepared by calcining a suspension of K₈Nb₆O₁₉ and ZnCl₂ in the presence of H₃BO₃ catalyst at 500 °C [50].

Solid-State with CuO Doping and Sol-Gel:

NiNb₂O₆ ceramics were produced by solid-state reaction and CuO doping, enabling sintering at lower temperatures while maintaining high densification and dielectric performance. The sol-gel method also produced fine NiNb₂O₆ nanoparticles at 700 °C, which demonstrated excellent properties upon further sintering [51].

Mechanochemical Synthesis:

Pure-phase MgNb₂O₆ was synthesized at 800 °C after 15 hours of high-energy ball milling, which reduced the necessary synthesis temperature by 100 °C compared to conventional methods [52].

Doped Solid-State Reaction:

Mn²⁺-doped Mg_{1-x}Mn_xTa₂O₆ (x = 0.02–0.12) ceramics were synthesized via a solid-state method, with the influence of Mn substitution on structural and dielectric properties thoroughly examined [53].

Conventional Solid-State Synthesis:

Polycrystalline BaNb₂O₆ was synthesized at temperatures above 1100 °C using the conventional solid-state reaction method. Increasing sintering time led to enhanced grain growth, higher density, and improved energy storage efficiency [54].

4. PROPERTIES OF AB₂O₆ COLUMBITE-TYPE OXIDES

The AB₂O₆ columbite-type oxides, particularly those based on vanadium and niobium, exhibit a wide range of intriguing electronic, optical, and thermoelectric properties, making them valuable candidates for various technological applications. Akhlaq Ahmed et al. (2024) have studied the thermoelectric properties of divanadate oxides such as MgV₂O₆, CaV₂O₆, and BaV₂O₆, reporting

band gaps of 3.20 eV, 2.14 eV, and 1.76 eV, respectively. The results obtained demonstrate the potential of these materials for thermoelectric power generation and their use in renewable energy systems [55].

Likely, Niobate-based oxides also exhibit semiconducting behavior, making them well suitable for high-voltage and high-temperature applications [56]. In this context, Kieran B. Spooner et al. (2021) explored the optical and thermoelectric properties of $BaBi_2O_6$, a columbite-type oxide, revealing a band gap of 2.6 eV and emphasizing its potential for thermoelectric applications due to its $PbSb_2O_6$ -type crystal structure [57]. Similarly, Basavaraju et al. (2021) investigated $MgNb_2O_6$ nanoparticles and found a comparatively large band gap of 4.4 eV, indicating the material's use for photocatalytic and sensor applications [58].

The optical properties of these compounds also attract significant attention. In 2023, José Fábio de Lima Nascimento et al. synthesized $CaNb_2O_6$ and obtained an optical band gap of 3.18 eV, further confirming the potential of niobium-based columbite-type oxides for optical and photocatalytic applications [59, 60]. The Raman and infrared vibrational modes of columbite-type compounds have been predicted using correlation methods [61] and further analyzed through theoretical calculations employing the Wilson GF matrix method [62]. Understanding of the electronic and optical properties of these materials has also been studied theoretically by using density functional theory (DFT). DFT calculations using the Wien2k package were utilized to improving the lattice constants and atomic positions in $CaNb_2O_6$, giving a direct band gap of 3.5 eV, making it suitable for photovoltaic applications. The calculations indicated that the valence band is mainly determined by Oxygen orbitals, while the conduction band is notably influence by both Calcium and Niobium, further increasing the material's electronic properties [63]. Furthermore, ab initio DFT simulations have been applied to MgV_2O_6 exploring its mechanical, structural, optical, electronic, and thermodynamic properties [64].

So we can conclude, the AB_2O_6 columbite-type oxides, especially Vanadates and Niobates, demonstrate a magnificent integration of semiconducting, optical, and thermoelectric properties, placing them as promising materials for many applications. Their capacity to conduct electricity under specific conditions, stability in high-temperature and high-voltage environments, makes them right choice for thermoelectric devices, photovoltaic cells, light-emitting diodes, and photocatalysts.

5. USES AND APPLICATIONS

Ternary metal oxides (TMOs), particularly ABO_4 and AB_2O_6 crystal structures, have recently gained significant attention in the field of gas sensing. These materials have a wide range of advantages including sensing capabilities, ease of synthesis, cost-effectiveness, and environmental friendliness. They offer considerable advantages in terms of material design adaptability and surface sensitivity also, being the crucial parameters in enhancing the selectivity and sensitivity of gas sensors, consequently TMOs play key role into next-generation sensors designed for environmental monitoring and industrial safety applications [65].

In the energy storage sector, Niobium-based ternary oxides have potential as anode materials for lithium-ion batteries. These materials, such as $MgNb_2O_6$, $CaNb_2O_6$, and $BaNb_2O_6$, show structural strength and outstanding electrochemical stability under high charge-discharge rates. They can undergo extended cycling without significant destruction makes them substitute to traditional graphite-based anodes, specifically for high-power applications such as electric vehicles and portable electronic devices [48].

There is rise in the use of cadmium-based phosphor materials, for optoelectronic devices particularly due to their efficiency and efficient photoluminescence. These materials are investigated in applications including solar cells, light-emitting diodes (LEDs), biosensors, and biomedical imaging. Considerably, cadmium selenide/zinc sulfide ($CdSe/ZnS$) core/shell quantum dots (QDs) are being researched for their role in cancer diagnostics and therapy. Although the inherent toxicity of cadmium imposes constraints for industrial and biomedical adoption, these QDs offer high brightness, reducing the X-ray dosage needed in medical imaging, thus reducing related health risks. Current research focuses on surface modification and encapsulation techniques to lessen their cytotoxic and genotoxic effects [66].

In the realm of photocatalysis and environmental remediation, $MgNb_2O_6$ nanoparticles have demonstrated exceptional photocatalytic activity under UV irradiation. These nanoparticles effectively degrade organic dyes such as Acid Red 88 (AR-88) and Fast Orange Red (FOR), indicating their potential for wastewater treatment applications. Additionally, $MgNb_2O_6$ nanoparticles show high sensitivity and electrochemical response in detecting paracetamol (acetaminophen), a commonly used pharmaceutical compound. Their responsive behavior over a wide potential range (-1.5 V to +0.3 V) positions them as strong candidates for

chemical sensing applications in pharmaceutical and environmental fields [57].

Microwave dielectric materials play a foundational role in the advancement of modern communication systems. ZnNb₂O₆ ceramics stand out as one of the most promising materials in this category, owing to their impressive dielectric constant ($\epsilon \sim 25$), high quality factor ($Q \times f = 83.7$ GHz), and a low temperature coefficient of resonant frequency ($\tau_f = -56$ ppm/°C). These properties make ZnNb₂O₆ an ideal material for the fabrication of dielectric resonators, filters, and antennas used in microwave communication devices. Furthermore, its relatively low sintering temperature (~ 1200 °C) makes it compatible with conventional ceramic processing techniques, thereby reducing manufacturing costs [67].

Cadmium-based quantum dots, especially CdSe/ZnSQDs, continue to be extensively researched for a wide range of high-performance applications, such as biomedical screening, solar energy harvesting, and LEDs. While cadmium's cytotoxic effects pose a challenge—particularly in clinical and therapeutic contexts—the remarkable optical properties of these QDs, including their tunable emission wavelengths and photostability, remain unmatched. The high brightness of cadmium-based phosphors enables reduced exposure times and lower doses in X-ray-based diagnostics, thereby offering a balance between performance and safety when used judiciously [68].

In photoelectrochemical (PEC) water splitting applications, NiV₂O₆ thin films have emerged as effective photoanode materials. Their synthesis and evaluation have revealed favorable electronic properties for solar-driven hydrogen production, a clean and sustainable alternative energy source. The efficient light absorption and charge transport characteristics of NiV₂O₆ make it a valuable component in PEC devices aimed at large-scale hydrogen generation [69].

ZnTa₂O₆ (ZTO) is a lead-free piezoelectric material that offers a combination of high thermal stability and excellent piezoelectric response. It crystallizes in the orthorhombic Pbcn space group and exhibits strong Raman activity. The material's dielectric constant (7.05), low loss tangent, moderate piezoelectric coefficient ($d_{33} = 20$ pC/N), and energy storage efficiency of 43% make it an attractive option for applications in energy harvesting, electronic actuators, and environmentally friendly transducers that operate under high-temperature conditions [70].

A novel electrocatalyst with the columbite–tantallite structure, Fe_{0.79}Mn_{0.21}Nb_{0.16}Ta_{0.84}O₆, has been developed for water splitting, demonstrating

efficient hydrogen evolution reaction (HER) and oxygen evolution reaction (OER) performance. The material delivers low overpotentials (190.2 mV for HER and 284.8 mV for OER) and exhibits fast reaction kinetics and long-term stability. Density functional theory (DFT) calculations support its outstanding catalytic activity by revealing low Gibbs free energy for intermediate reaction steps and minimal OER overpotential, confirming its suitability for renewable hydrogen production [71].

Finally, an advanced electrochemical gas sensor based on a stabilized zirconia platform and using MnNb₂O₆ as the sensing electrode (SE) is fabricated to detect trace levels of sulfur dioxide (SO₂) at elevated temperatures. Among various electrode sintering conditions, the MnNb₂O₆-SE calcined at 1000 °C delivered the best performance, offering a strong voltage response (-27 mV to 5 ppm SO₂), a notably low detection limit (50 ppb), and a swift response time (37 seconds for 1 ppm SO₂ at 700 °C). This sensor also displayed outstanding linearity, stability, and selectivity against interfering gases, proving it a reliable tool for real-time gas monitoring in extreme environments. The detection mechanism was linked to a mixed potential model, where both ionic and electronic contributions govern sensor output [72].

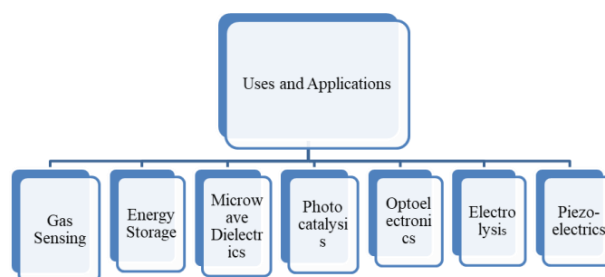


Figure 2. Uses and Applications of columbite-type oxides

6. FUTURE POTENTIALS

The future potential of Columbite Oxides and related ternary oxides is immense, spanning numerous high-impact fields such as energy storage, optoelectronics, photocatalysis, and biological applications. NiV₂O₆ is one of the important Columbite Oxides with immense future potential given as below:

NiV₂O₆, with its direct band gap of 0.172 eV, has the potential as a semiconductor that could be employed for thermal shielding and optoelectronic devices. The versatility of its electronic properties could open the door for its incorporation on into advanced optoelectronic devices, such as photodetectors, modulators, and lasers, offering efficient performance in energy conversion technologies [73].

Furthermore, NiV_2O_6 has demonstrated exceptional photocatalytic activity, particularly when integrated into nanostructures like the Eu^{3+} -doped $MgNb_2O_6$ nanophosphors. These materials exhibited significant photocatalytic degradation of organic dyes like AR-88, as evidenced by the Langmuir–Hinshelwood model, showing great potential for environmental remediation applications, such as wastewater treatment, and elevating photocatalytic processes for energy harvesting. As such, NiV_2O_6 could be investigated for use in sustainable energy applications, particularly in the field of photocatalysis, where it could promote both the degradation of pollutants and the synthesis of renewable energy sources [74,75].

The potential of NiV_2O_6 in the field of biological applications is also promising. The nanoengineered NiV_2O_6 nanoflowers exhibited enhanced oxidase-like activity, boosting fluorescence by a remarkable 8-fold. This increase in fluorescence was attributed to the unique properties of the nanostructures, suggesting their application in biosensing. Through radical scavenger studies, insights into the underlying mechanism of this enhancement were gathered, and the material's sensitivity was demonstrated in detecting low concentrations of glutathione (0.05–12.5 μM) in serum and cancer cells. This highlights its potential as a powerful bioanalytical tool, especially in diagnostic applications and cancer therapy monitoring [76].

The integration of NiV_2O_6 into energy storage technologies also holds significant promise. The graphene/ MnV_2O_6 nanocomposite demonstrated excellent electrochemical performance, making it a strong contender for high-efficiency electrode materials in supercapacitor applications. The composite's enhanced charge storage capabilities could contribute to the development of advanced energy storage devices with higher energy density and faster charge-discharge cycles, essential for next-generation electronics, electric vehicles, and portable devices [77].

Additionally, the DFT-based analysis of MgV_2O_6 confirmed its mechanical stability and semiconducting nature, with a band gap of 2.195 eV. The material's optical properties suggest its suitability for solar heat shielding applications, where it could play a role in passive energy management systems for buildings and solar energy devices, contributing to both energy conservation and efficiency improvements [60].

The future applications of NiV_2O_6 and related ternary metal oxides extend across a wide range of high-tech domains, from energy storage to environmental sustainability and medical

diagnostics. The material's unique properties, such as its band gap, fluorescence enhancement, and photocatalytic capabilities, position it as a versatile material for advanced technologies, offering vast potential for innovation in both industrial and biomedical fields. As research progresses, it is expected that these materials will undergo further optimization to unlock even more applications, driving the development of more efficient, sustainable, and high-performance technologies [54,60,73-77].

In recent years, AB_2O_6 oxide based phosphor materials with columbite type structure have attracted increasing attention due to their promising photoluminescence properties as well as potential forensic applications. In this context, a recent study focusing on $CdNb_2O_6$ [78] investigated both the luminescence behavior and latent fingerprint detection capabilities of this host material. The results demonstrated strong visible emission, indicating that $CdNb_2O_6$ can serve as an efficient phosphor when doped with appropriate activator ions. Moreover, its surface morphology and optical characteristics enabled high-contrast visualization of latent fingerprints under UV excitation, emphasizing its suitability for forensic imaging. This work highlights the dual-functional nature of columbite-type hosts like $CdNb_2O_6$, which are not only viable for lighting and display technologies but also hold significant promise in the field of forensic science, particularly in advanced latent fingerprint visualization techniques.

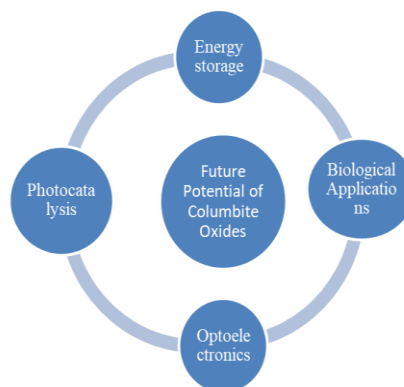


Figure 3: Future Potential of Columbite Oxides

7. CONCLUSION

Columbite-type AB_2O_6 oxides hold great promise for future technology. Their distinctive crystal structures and highly flexible synthesis approaches enable scientists to improve their properties to suit diverse applications. The materials are being used for exciting uses like converting heat into electricity, purifying pollution with light, enhancing battery storage, and upgrading electronic devices. Large-scale applications, however, are not simple. They usually need extremely high temperatures to synthesize,

are poor absorbers of sunlight, are poor conductors of electricity, and rely on expensive or hard-to-get rare elements. To get past these issues, scientists have to continue to develop new methods for enhancing them, like chemical adjustments or computational simulations to forecast how they will act. In the future, through the improvement of their synthesis and use, these oxides may be used as an integral component of greener, smarter, and more efficient technologies. They have the potential to drive everything from environmentally friendly electronics to clean energy solutions.

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IZVOD

KOLUMBITSKI STRUKTURIRANI AB₂O₆ OKSIDI: RAZUMEVANJE NJIHOVE STRUKTURE, SINTEZE I POTENCIJALA ZA BUDUĆE TEHNOLOGIJE

Oksidi AB₂O₆ su nedavno privukli značajnu pažnju zbog svojih ekskluzivnih svojstava. Ova jedinjenja pokazuju raznovrsne strukturne oblike sa različitim karakteristikama. Među njima, oksidi kolumbitskog tipa poseduju ortorombsku strukturu, pripadaju prostornoj grupi Pbcn i pokazuju simetriju D_{2h}¹⁴. Struktura ovog materijala sastoji se od oktaedara AO₆ i BO₆ koji su raspoređeni cik-cak i imaju snažan uticaj na karakteristike ovih materijala. Ovi materijali igraju vitalnu ulogu u različitim primenama kao što su mikrotalasna tehnologija, kataliza, skladištenje energije, elektronski senzori i optički materijali. Na osnovu sveobuhvatnog pregleda literature, ovaj pregled je obuhvatio detaljnu studiju strukture, fizičkih svojstava, tehnika sinteze i primene ovih oksida. Nakon toga, razmatramo budući potencijal kolumbitskih oksida, sa naglaskom na strategije i računarsko modeliranje za poboljšanje njihovih svojstava i performansi za buduće tehnologije.

Ključne reči: kolumbit, termoelektrika, mikrotalasna tehnologija, računarsko modeliranje

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Rachana Gaur:	https://orcid.org/0009-0007-8862-6903
Dr. Ruby Jindal:	https://orcid.org/0000-0002-8589-5549
Archana Tripathi:	https://orcid.org/0000-0002-1811-284X
Harleen Kaur:	https://orcid.org/0000-0002-5034-2119
Aparna Shekhar:	https://orcid.org/0000-0003-2286-2587