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A Review of Computational and Theoretical Studies on Anti-Perovskite Solids

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Abstract

Anti-perovskite materials are really turning heads lately, and it's no wonder why. Their unique structural inversion, unlike traditional perovskites, leads to some truly remarkable properties in terms of electronics, thermal behavior, and magnetism. With a general formula of A₃BX, these materials are known for boasting high ionic conductivity and less thermal conductivity; they show potential for topological and superconducting behaviors. This makes them perfect candidates for all sorts of applications, from solid-state batteries to thermoelectrics, spintronics, and even quantum computing. Researchers have made great strides using computational and theoretical studies especially with density functional theory (DFT) to predict how stable these structures are, along with their electronic configurations and overall functional properties. High-throughput screening, phonon dispersion analysis, and machine learning techniques are also helping push the discovery of new anti-perovskite compounds along faster than ever. Yet, there are still hurdles to overcome, like accurately modeling electron correlation effects and lattice anharmonicity, not to mention the limited experimental validation. Still, the collaboration between computational predictions and experimental work opens up exciting possibilities for designing tailored anti-perovskites. With their multifunctional traits, they really could be game-changers for future energy and electronic technologies.

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Introduction

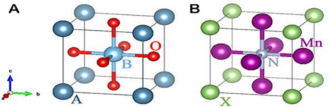
Recently, anti-perovskite materials have started to gain traction as exciting options for a variety of technological advancements. Their unique structural, electronic, magnetic, and thermal properties set them apart. Unlike traditional perovskites, anti-perovskites feature a flipped crystal arrangement, where the anions and cations switch places. brings about inversion innovative electronic configurations and results in remarkable material characteristics, such as impressive ionic conductivity, low thermal conductivity, and the potential for superconductivity (Zhang et al., 2017). These features make anti-perovskites particularly promising for cutting-edge devices like solidbatteries. thermoelectrics. and Computational and theoretical approaches are crucial for understanding and designing anti-perovskite materials. Techniques like density functional theory (DFT) and other first-principles methods are widely used to delve into aspects such as structural stability, electronic band structures, phonon dispersions, and defect characteristics of these materials. For instance, Sun et al. (2015) utilized DFT to investigate the thermodynamic stability and electronic properties of antiperovskite lithium-rich compounds, highlighting their promise as solid electrolytes for lithium-ion batteries. Likewise, research on transition metal-based anti-perovskites has unveiled intricate magnetic phase diagrams and intriguing topological features, suggesting their potential applications in quantum computing and magnetocaloric technologies, as noted by (Kumar & Schwingenschlögl, 2017). One of the biggest perks of using computational

studies is that they can predict new compounds and their stability phases without having to rush into experimental synthesis. By creating phase diagrams and running molecular dynamics simulations, researchers can take a closer look at how temperature and pressure affect material behaviors and pinpoint the best conditions for stability (Gao et al., 2018). This theoretical understanding really cuts down on the trialand-error typically involved in experimental work, helping to speed up the discovery of innovative functional materials. Computational models have really opened our eyes to the tolerance factors, lattice dynamics, and defect chemistry that play a huge role in the stability and performance of antiperovskites. For instance, the ideas around geometric and electronic tolerance factors help us predict how likely it is to create stable anti-perovskite phases (Filippetti & Fiorentini, 2019). Getting a grip on these parameters is key to finetuning properties like ionic conductivity and magnetic ordering, which are quite sensitive to changes in structure and chemical substitutions. While we've made some impressive strides, we're still facing hurdles when it comes to precisely modeling strong electron correlation effects and the anharmonic behavior of lattices in certain anti-perovskite systems. Luckily, the ongoing advancements in hybrid functionals, GW approximations, and even machine learning techniques show a lot of potential for tackling these challenges and enhancing the predictive capabilities of our computational models (Liu et al., 2020).

Structural Characteristics Of Anti – Perovskites Antiperovskites are fascinating materials that feature an unusual

arrangement of ions, setting them apart from the more common perovskites. In a traditional perovskite, the structure follows the formula ABX3, where A and B are cations and X is usually an anion like oxygen. However, anti-perovskites use the A3BX formula instead. In this configuration, the larger metal cations (A) create a cubic lattice, while the smaller non-metal or metalloid atom (B) sits in the center of the octahedra, and the anion (X) is located at the center of the cube. This inversion in positioning results in significantly different structural and bonding properties compared to their conventional perovskite counterparts (Zhang *et al.*, 2017).

The typical crystal structure you'll find in anti-perovskites is the cubic Pm-3m space group. In this setup, the A-site cations create a face-centered cubic (FCC) lattice. Meanwhile, the B atom usually one of the elements like nitrogen, oxygen, or carbon occupies the body center, nicely enclosed by a cage formed by the A-site atoms. This arrangement leads to robust A–B and A–X interactions, which play a pivotal role in shaping the electronic, magnetic, and ionic properties (Filippetti & Fiorentini, 2019).



One of the standout characteristics of anti-perovskites is their impressive structural flexibility. This flexibility opens the door for significant substitutions at both the A and B sites. Being able to tweak these elements is vital for customizing the material properties, such as ionic conductivity and magnetism.

Computational Methods for Anti-Perovskite Discovery

Computational techniques play a crucial role in predicting, designing, and optimizing anti-perovskite materials. Here are some commonly used methods and approaches:

A.Density Functional Theory (DFT):

The first-principles method most often used to explore the electronic structure, phase stability, magnetic properties, and mechanical behavior of anti-perovskites. It's a great tool for pinpointing stable crystal structures and predicting things like band gaps, phonon spectra, and elastic properties. Sun, Zhang, & Ceder (2015)

B.High-Throughput Computational Screening:

Automated DFT calculations are used to screen large material databases for stable anti-perovskite candidates. Properties such as formation energy, structural stability, and ionic conductivity are rapidly evaluated. Zhang, Zhao, & Ceder (2017)

C.Phonon Dispersion and Thermodynamic Stability Analysis:

Phonon calculations help us understand how stable certain structures are by looking for those pesky imaginary frequencies. On the other hand, thermodynamic models take a closer look at Gibbs free energy as conditions change with temperature and pressure. Gao *et al.* (2018)

D.Machine Learning Models:

This approach uses DFT datasets to forecast important structural and functional properties, including stability, conductivity, and magnetism. It significantly accelerates and broadens the material discovery process, surpassing the capabilities of traditional DFT methods on their own. Liu, Guo, & Wang (2020)

Prediction and Design of Novel Anti-Provskites:

Anti-perovskite materials are heavily dependent on sophisticated computational methods that empower researchers to efficiently and accurately sift through a vast array of compositional possibilities. A key emphasis is on pinpointing compositions that not only demonstrate thermodynamic stability but also possess attractive electronic characteristics and are structurally viable. Density Functional Theory (DFT) is essential in forecasting the formation energies, electronic band structures, and magnetic ordering of anti-perovskite compounds (Sun et al., 2015). Highthroughput computational screening has become a go-to method for predicting new stable A₃BX-type anti-perovskites by sifting through vast databases of elemental combinations. These techniques examine crucial factors such as tolerance factors, lattice constants, and cohesive energies to assess structural stability (Zhang et al., 2017). Moreover, phonon dispersion analysis is utilized to confirm dynamic stability by identifying the absence of any imaginary vibrational modes (Gao et al., 2018). Machine learning (ML) algorithms are becoming more and more integrated with quantum mechanical simulations, paving the way for faster discoveries in material science. By utilizing ML models that are trained on existing anti-perovskite datasets, researchers can effectively predict the stability and functional properties of new, untested compounds. This not only speeds up the process but also cuts down on the computational costs (Liu et al., 2020). With these innovative methods, it's now possible to quickly pinpoint candidate materials suitable for use in solid-state batteries, thermoelectrics, and spintronics. Ultimately, bringing together computational modeling, materials informatics and data-driven techniques has truly changed the game in designing new anti-perovskites. This approach opens up exciting possibilities for uncovering materials with specific properties that can be used in cuttingedge technologies.

Electronic and Physical Properties

Anti-perovskite materials boast a fascinating range of electronic and physical properties that make them exceptionally appealing for uses in energy storage, electronics, thermoelectrics, and quantum technologies. These unique traits stem from their distinctive crystal structure and the ability to mix and match various elements at different lattice sites.

Electronic Properties:

•Band Structure and Conductivity:

Anti-perovskites display a fascinating variety of electronic characteristics, ranging from metallic to semiconducting, based largely on their chemical makeup. Take, for example, transition-metal-based anti-perovskites like Mn₃GaN and Mn₃Sn; these materials exhibit semi-metallic properties along with significant spin-polarization, making them excellent candidates for spintronic devices (Kumar Schwingenschlögl, 2017). On the other hand, some lithiumrich anti-perovskites may act as electronic insulators, yet they shine in the realm of ionic conductivity, which is perfect for solid-state electrolytes used in lithium-ion batteries (Sun et al., 2015).

•Topological Characteristics:

Certain anti-perovskites are expected to showcase fascinating topological states like Dirac or Weyl semimetals. This has caught the attention of researchers diving into quantum computing and low-power electronics (Zhang *et al.*, 2017).

Physical Properties:

•Magnetic Behavior:-Transition metal anti-perovskites have a fascinating ability to show complex magnetic behaviors, including antiferromagnetism and non-collinear spin structures. This characteristic positions them as strong candidates for applications in magnetocaloric technology and spintronics (Kumar & Schwingenschlögl, 2017).

•Thermal Conductivity:

Many anti-perovskite materials have incredibly low thermal conductivity thanks to their flexible lattice dynamics, which makes them quite promising for use in thermoelectric devices (Gao *et al.*, 2018).

•Mechanical and Structural Flexibility:

The cubic symmetry along with its capacity to handle substitutions at various sites gives it impressive mechanical strength and the ability to adjust physical characteristics, such as elastic moduli and thermal expansion (Filippetti & Fiorentini, 2019).

Anti-perovskites are really fascinating materials that offer a wide range of electronic and physical properties that can be customized. By tweaking their chemical composition and using computational modeling, we can fine-tune their characteristics, making them incredibly adaptable for the technologies of the future.

Challenges and Limitations

Accuracy of Standard DFT Methods

A lot of computational studies lean on Density Functional Theory (DFT), but it can fall short when dealing with systems that have strongly correlated electrons, particularly in transition-metal-based anti-perovskites. Standard exchange-correlation functionals like GGA or LDA tend to underestimate band gaps and often miss the mark on accurately capturing magnetic interactions (Liu *et al.*, 2020)

•Computational Cost of Advanced Methods

While more precise methods like hybrid functionals (such as HSE06), DFT+U, or GW approximations yield superior outcomes, they demand a lot of computational power. This makes them less suitable for high-throughput screening (Filippetti & Fiorentini, 2019).

•Modeling Dynamic and Anharmonic Effects

Anti-perovskite materials tend to show lattice anharmonicity and soft phonon modes, which means that most harmonic approximations in phonon calculations can't capture the full picture. This limitation can affect the accuracy of predictions regarding vibrational and thermal stability (Gao *et al.*, 2018).

•Limited Experimental Benchmarking

It's challenging to test out theoretical predictions about antiperovskites since many of them haven't been synthesized yet, making it tough to validate the computational forecasts and improve our models (Zhang *et al.*, 2017).

Applications of Anti- Perovskites:

Anti-perovskite materials have captured a lot of interest lately, and it's easy to see why. Their amazing structural flexibility and multifunctional properties open the door to a host of applications in some really innovative technologies.

I.Solid-State Electrolytes in Batteries

Anti-perovskites like Li₃OCl and Li₃OBr have demonstrated impressive lithium-ion conductivity, positioning them as exciting candidates for solid electrolytes in all-solid-state lithium-ion batteries. Their low activation energy and rapid ion transport not only bolster battery safety but also enhance overall performance (Zhu *et al.*, 2015; Sun *et al.*, 2015).

II.Thermoelectric Devices

Some anti-perovskite materials show not only low thermal conductivity but also adjustable electrical conductivity traits that make them pretty appealing for thermoelectric applications. For instance, compounds like Mn₃AN (where A can be Cu, Zn, or Sn) have displayed encouraging thermoelectric performance, as noted by (Zhang *et al.*, 2017).

III.Spintronics and Magnetic Devices

Transition metal anti-perovskites, like Mn₃Sn and Mn₃Ga, exhibit unique non-collinear magnetic ordering along with impressive anomalous Hall effects. These characteristics make them quite useful in the realm of spintronic devices, particularly for applications in memory storage and magnetic sensing (Kumar & Schwingenschlögl, 2017).

IV.Topological Quantum Materials

Anti-perovskites are thought to showcase some fascinating properties, such as topologically non-trivial states that include behaviors like Dirac and Weyl semimetals. These characteristics hold exciting potential for innovations in quantum computing and advanced electronic devices (Liu *et al.*, 2020).

Future Prospects

The future looks incredibly bright for anti-perovskite materials! Thanks to their adjustable structures, versatile properties, and ability to work with various cutting-edge technologies, they're getting a lot of attention. As research progresses, we can expect some really exciting pathways that will drive the development and use of these materials. We're witnessing some exciting advancements in computational methods these days, especially with machine learning and high-throughput DFT simulations. These innovations are set to speed up the discovery of new anti-perovskite compositions. Thanks to these powerful tools, we can now predict crucial properties like stability, ionic conductivity, and magnetism over a wide range of chemical spaces, which cuts down the need for extensive experimental testing (Liu et al., 2020). Next up, the evolution of cutting-edge solid-state batteries stands to gain a lot from anti-perovskite-based electrolytes. This is particularly true as we uncover new compositions that boast better ionic conductivity and chemical stability (Sun et al., 2015). Grasping the stability of interfaces and exploring various doping strategies will be key in this area. Looking further, the topological and quantum applications of anti-perovskites are set to take off, especially with their potential in spintronic devices and quantum computers. Some anti-perovskites are even expected to show Dirac and Weyl semimetal characteristics, which are vital for efficient electronic transport with minimal dissipation (Zhang et al., 2017). To truly harness the potential of metastable anti-perovskite phases and nanostructures that theories suggest, we need to see some growth in experimental synthesis techniques. It's essential that we align theoretical modeling with practical advancements. This integration will be crucial for unlocking the vast opportunities that antiperovskites offer in areas like energy storage, electronics, and quantum technology.

Conclusion

Anti-perovskite materials are gaining a lot of attention as an exciting new category of compounds, thanks to their unique crystal structure and adaptability. Their ability to flexibly change structure, combined with fascinating electronic, thermal, and magnetic properties, positions them perfectly for cutting-edge technologies like solid-state batteries, thermoelectric devices, spintronic components, and quantum materials. What sets anti-perovskites apart from traditional

perovskites is their capacity for broad chemical substitutions at various sites. This flexibility allows scientists to fine-tune key characteristics like ionic conductivity, magnetism, and topological behaviors. Thanks to computational methods particularly density functional theory (DFT) and highthroughput screening—the discovery and design innovative anti-perovskite phases have been sped up significantly. These techniques aid in predicting important aspects such as phase stability, band structure, phonon behavior, and defect chemistry, all without the need for immediate experimental synthesis. Moreover, incorporation of machine learning into this process is boosting both accuracy and efficiency. That said, there are Reference

- Filippetti, A., & Fiorentini, V. (2019). Tolerance factors in perovskite and anti-perovskite materials. Journal of Materials Chemistry A, 7(3), 1048–1056. https://doi.org/10.1039/C8TA10323G
- Gao, T., Han, D., Zhu, Y., & Wang, D. (2018). First-principles phonon and stability analysis of anti-perovskites. Physical Chemistry Chemical Physics, 20(45), 28567–28574. https://doi.org/10.1039/C8CP05341E
- Gao, T., Han, D., Zhu, Y., & Wang, D. (2018). Theoretical prediction of temperature-dependent phase stability in anti-perovskite compounds. Physical Chemistry Chemical Physics, 20(45), 28567–28574. https://doi.org/10.1039/C8CP05341E
- Kumar, A., & Schwingenschlögl, U. (2017). Magnetic and topological properties of anti-perovskites: A DFT study. Chemistry of Materials, 29(10), 4200–4206. https://doi.org/10.1021/acs.chemmater.7b01145
- Liu, J., Guo, Y., & Wang, Y. (2020). Machine learning assisted prediction of anti-perovskite material

still some hurdles to overcome. These include challenges in modeling strongly correlated electrons, dealing with anharmonic lattice vibrations, and a lack of experimental validation for some predicted materials. Plus, many antiperovskite compounds tend to be metastable or tricky to synthesize. Despite these challenges, the outlook for antiperovskites is promising. Ongoing progress in computational modeling, materials informatics, and experimental techniques are vital for tapping into their full potential. With collaborative research efforts, anti-perovskites could soon change the game in fields such as energy storage, electronics, and quantum materials.

- properties. npj Computational Materials, 6, 45. https://doi.org/10.1038/s41524-020-0313-7
- Sun, Y., Zhang, Y., & Ceder, G. (2015). First-principles investigation of anti-perovskite lithium conductors. Chemistry of Materials, 27(20), 7443–7450. https://doi.org/10.1021/acs.chemmater.5b03260
- Zhang, Y., Zhao, Y., & Ceder, G. (2017). Computational design of functional anti-perovskites. Nature Reviews Materials, 2, 17023. https://doi.org/10.1038/natrevmats.2017.23
- Zhang, Y., Zhao, Y., & Ceder, G. (2017). Computational design of functional anti-perovskites. Nature Reviews Materials, 2, 17023. https://doi.org/10.1038/natrevmats.2017.23
- Zhu, Y., He, X., & Mo, Y. (2015). ACS Applied Materials & Interfaces, 7(42), 23685–23693. https://doi.org/10.1021/acsami.5b07517