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Modulation on terahertz absorption properties in Ln^{III}–[Ag^I(CN)₂]₃ networks†

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Molecular materials are playing a pivotal role in the advancement of prospective THz technologies. Herein, we have prepared nine isostructural coordination complexes of general formula [Ln^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**LnAg**, Ln = La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, and Dy). Vibrational spectroscopy has unveiled gradual changes in the $\nu(\text{C}\equiv\text{N})$ stretching frequencies within both infrared (IR) and Raman spectra from **LaAg** to **DyAg**, stemming from incremental alterations in crystallographic unit cell parameters and volumes. Similarly, the THz spectra manifest pronounced absorption signals between 1.0 to 1.2 THz, which can be judiciously tuned by varying lanthanide(III) ion and measurement temperature. *Ab initio* density-functional theory (DFT) calculations of THz spectra elucidate the primary contribution from opposite translational vibrations of Ag atoms and O atoms from H₂O around Ln(III) atoms. Furthermore, thin films of **LnAg** can achieve a large return loss in a broad bandwidth in impedance-matching simulations.

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Introduction

Research on terahertz (THz) radiation, spanning from 100 gigahertz (GHz) to 30 THz between microwaves and infrared lights and is essentially terra incognita, has recently garnered increasing attention from scientists all over the world.^{1–3} Technological innovations in laser technology, nanotechnology, and photonics are now making the previously unexploited THz frequency band accessible in many fields.^{4–8} In particular, THz time-domain spectroscopy (THz-TDS) is emerging as a cutting-edge method to analyze a wide variety of materials with valuable insights into the fingerprint of the molecular structures, building block arrangements, and inter-

intramolecular interactions.^{9–13} Combining THz spectroscopy with a magnetic field enables the precise characterization of zero-field splitting, providing detailed insights into the electronic structure and magnetic properties of paramagnetic systems.^{14–16} THz light exhibits versatility in diverse applications, including non-invasive medical imaging and cancer diagnosis, material characterization, monitoring of toxic chemical compounds, sensing events due to natural and human-related disasters, and catalysis.^{4,17–20} Moreover, the THz band offers unprecedented channel capacities and is poised to contribute to the establishment of high bandwidth wireless sixth-generation (6G) networking systems with anticipated data rates reaching Tbit per s in the near future.^{21–24}

As society enters the Internet of Things (IoT) era, the pervasive expansion of wireless technology has introduced novel challenges, concerning excessive electromagnetic (EM) radiation pollution, EM interference, and EM compatibility issues.^{25–27} Additionally, the manipulation of THz radiation tailored for diverse communication systems is still in its infancy and is a subject of active research.^{22,23} Hence, there is a compelling imperative to develop new materials capable of achieving efficient THz absorbers while affording THz tunability to modulate different frequency bands on demand. Particularly, molecular complexes are emerging as an appealing category of materials owing to their simple and adjustable structures and predisposition to merge various functionalities, such as magnetism, luminescence, non-linear optics, and low-frequency absorbing capability.^{28–36} Moreover, the desired THz

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† Electronic supplementary information (ESI) available: Legends for supporting movie; single crystal X-ray diffraction analyses; thermogravimetric data; powder X-ray diffraction studies; IR absorption and Raman scattering spectroscopies; terahertz time-domain spectroscopy; first-principles phonon mode calculation data. CCDC 2339311 (**LaAg**) and 2339312 (**DyAg**). For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d4qi00800f>



absorbing capability can be introduced by meticulously choosing building blocks and arranging them in solid-state structures. This strategic approach harnesses the inherent flexibility of molecular complexes, facilitating the deliberate engineering of materials with tailored THz characteristics.

Up to now, a diverse array of versatile materials based on metal complexes have been designed and prepared.^{37–41} Among them, lanthanide ions (Ln(III)) stand out as one of the most promising metal centers for the assembly of diverse molecular complexes.^{42–47} Notably, similarity in geometry and charges together with the lanthanide contraction offer lanthanide ions as promising candidates to achieve controllable tuning effects on structures and various properties. For instance, Wong *et al.* reported the study of second harmonic generation (SHG) as well as quadratic hyperpolarizability for a series of compounds $[(C_9H_7O_2)_3Ln]_n$, in which both non-linear optical (NLO) properties feature a similar dependence over the lanthanide series with maxima of SHG for Gd(III) and hyperpolarizability for Tb(III).⁴⁸ Besides, it was reported $[Ln(dma)_5][W(CN)_8]$ compounds exhibited diverse SHG properties with SH susceptibilities ranging from 4.7×10^{-12} to 9.4×10^{-11} esu.⁴⁹ Furthermore, a smooth and continuous increase for both the C≡N stretching and bending modes in the Raman spectroscopy was reported for a sequence of samples $[Ln(H_2O)_3][Au(CN)_2]_3$ from larger Ln³⁺ ionic radius to smaller one, signifying great tuning effects *via* changing lanthanides.⁵⁰ However, compared to the extensive studies for magnetic, luminescence, spectroscopic, and NLO properties for Ln(III) complexes, the lanthanide tuning effects have been rarely studied in the THz region.^{48–54} Concurrently, silver ions (Ag(I)) compounds have appeared to be noteworthy candidates to create THz absorbing materials. Notably, the phonon spectra of AgCN compound exhibit low-frequency THz peaks at *ca.* 4 meV (0.96 THz), and various Ag(I) complexes have been observed to manifest broad terahertz absorbing behaviors within the range of 0.2–3 THz.^{55,56}

Therefore, in the present study, we prepared a series of $[Ln^{III}(H_2O)_3][Ag^I(CN)_2]_3$ (LnAg, Ln = La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, and Dy) complexes and investigated their tunable vibrational spectra and temperature-dependent THz absorption properties. Furthermore, we conducted density-functional theory (DFT) calculations of the THz spectra and performed impedance-matching simulations to validate the observed phenomena and potential applications.

Results and discussion

Structural studies

The crystallographic data of two representative crystals LnAg and DyAg, obtained as a result of a slow evaporation of the diluted aqueous solution containing $LnCl_3$ (Ln = La(III) and Dy(III)) and $K[Ag(CN)_2]$, were collected at *ca.* 90 K in the single crystal X-ray diffraction measurement (refer to the details in the Experimental section, Table S1†). Both compounds are isostructural and crystallize in a three-dimensional (3D) network

with a hexagonal $P6_3/mcm$ space group. Analogously, the isostructural nature with the same space group aligns with observations in other reported dicyanoaurate and dicyanoargentate complexes in the literature.^{34,50,57–60} Nevertheless, the unit cell parameters (*a*, *b*, and *c*) and volumes (*V*) of DyAg are smaller than LnAg with a shrinkage of 1.4%, 1.4%, 3.0%, and 5.8% in *a*, *b*, *c*, and *V*, respectively, arising from the lanthanide contraction.^{61–63} The lanthanum(III) and dysprosium(III) ions are nine coordinates, including six nitrogen atoms originating from the cyanide ligand arranged in prismatic geometry and three coplanar oxygen atoms from the water (Fig. 1a). The average bond distances of $\langle Ln-O \rangle$ are 2.529(9) and 2.414(7) Å for LnAg and DyAg, respectively, slightly shorter than 2.653(7) Å for $\langle La-N \rangle$ and 2.505(5) Å for $\langle Dy-N \rangle$ (Table S2†). The linear configuration of $[Ag(CN)_2]^-$ ions with average bond angles $\langle Ag-C-N \rangle$ of 178.7(8)° and 179.8(8)° for LnAg and DyAg, respectively, connects with two adjacent Ln(III) ions along the *c*-axis (Fig. 1b and Fig. S1, Table S3†).

Each Ag atom is arranged with four near Ag atoms in the *ab*-plane, thereby forming Ag–Ag interactions with typical distances of 3.3241(3) Å for LnAg and 3.2762(8) Å for DyAg (Table S2†). Such Ag–Ag interaction distances are comparable with the other isostructural complexes with different lanthanide ions.^{50,57,58} The 3D networks with argentophilic interactions as well as hydrogen-bonding interactions between the water and nitrogen atoms of cyanide lead to relatively robust thermal stability until 100 °C. Further heating results in the dehydration of three water molecules above 100 °C and the decomposition of cyanides exceeding *ca.* 200 °C (Fig. S2†).

The phase purity and isostructural nature of all nine single crystals and the powdered LnAg samples were confirmed by the good agreement between the experimental powder X-ray diffraction (PXRD) patterns and the simulated one based on the results of single-crystal X-ray analysis of the LnAg crystal structure (Fig. 1c). Moreover, the gradual shift of the peak position to higher 2θ angles is observed for three selected peaks with Miller indices (002), (102), and (004) from LnAg to DyAg (Fig. 1d and S3†), which agrees with the decreased unit cells, especially along *c*-axis, and volumes from crystallographic data. The tunable structural parameters concerning different Ln(III) ions open up possibilities of finely modulating the structure-related vibrational properties, especially in the rarely explored THz region, which depends on the crystal packing.

Vibrational properties

The infrared (IR) absorption and Raman scattering spectra were further investigated to elucidate the gradual transitions among diverse LnAg samples. The prominent features in the room-temperature IR spectra of LnAg comprise one principal $\nu(C\equiv N)$ stretching band between 2130 to 2170 cm^{-1} , $\nu(O-H)$ symmetric and asymmetric stretching bands at *ca.* 3600 and 3534 cm^{-1} , respectively, as well as $\delta(H-O-H)$ bending band at around 1609 cm^{-1} (Fig. 1a and S5†). In parallel, the Raman spectra of LnAg are characterized by one intense $\nu(C\equiv N)$ stretching band between 2150 to 2170 cm^{-1} and several bending modes between 250 to 450 cm^{-1} , and weak Raman-



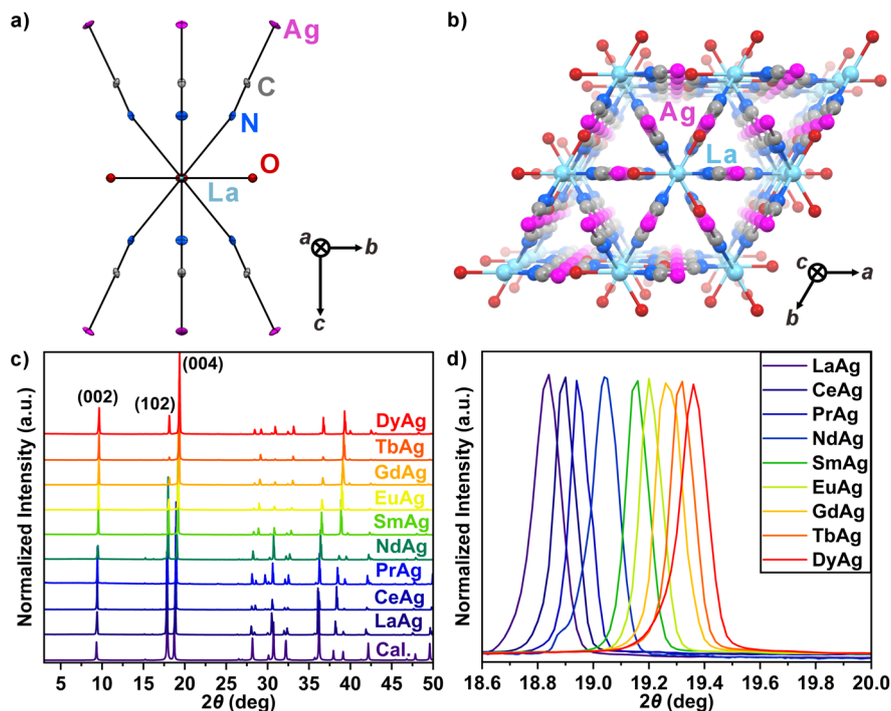


Fig. 1 (a) Representative structural unit of **LaAg** with atoms labeling along the *a*-axis measured at 90 K. Thermal ellipsoids of 50% probability are shown. (b) Crystal packing of **LaAg** along the *c*-axis. Color: La, cyan; Ag, pink; C, dark gray; N, blue; O, red. All hydrogens are omitted for clarity. (c) Experimental powder X-ray diffraction (PXRD) patterns of different **LnAg** samples compared with the calculated (Cal.) one based on the single-crystal structure of **LaAg** at 90 K after normalization. The Miller indices are marked for three indicated peaks. (d) Normalized PXRD patterns for (004) Miller index of different **LnAg** samples in the zoomed region.

active $\nu(\text{O-H})$ stretching bands at *ca.* 3540 cm^{-1} (Fig. 1b and $S6^\dagger$). Some extra broad and intensive peaks are unexpectedly observed for **EuAg** ($3260, 2088, 1913,$ and 947 cm^{-1}) and **SmAg** ($2535, 1913, 1703, 1673,$ and 818 cm^{-1}), consistently observed across different positions of crystals and experimental settings. Such phenomenon can be explained by the fluorescence interference in Raman spectroscopy utilizing 532 nm lasers, which appears to correspond to the excitation process of **EuAg** and **SmAg** selectively to a higher electronic state and then relaxed to generate fluorescence light.^{34,64,65} The relatively inefficient Raman scattering can be easily masked by the fluorescence process, thereby leading to relatively weak $\nu(\text{C}\equiv\text{N})$ and $\nu(\text{O-H})$ signals.

Remarkably, a discernible and gradual shift in $\nu(\text{C}\equiv\text{N})$ stretching frequencies for both IR and Raman spectra is observed from **LaAg** to **DyAg** (Fig. 2a and b). The IR-active $\nu(\text{C}\equiv\text{N})$ stretching vibrations have slightly lower energies, ranging from 2150 to 2157 cm^{-1} , whereas the Raman $\nu(\text{C}\equiv\text{N})$ modes span from 2159 to 2163 cm^{-1} . This shift demonstrates a linear correlation with the lanthanide ionic radii, within experimental error and setup resolution (Fig. 2c), akin to previously reported dicyanoaurate studies.⁵⁸ The shift in the $\nu(\text{C}\equiv\text{N})$ frequency associated with different **LnAg** samples is more obvious in IR spectra than in Raman spectra. In general, modifications in $\nu(\text{C}\equiv\text{N})$ stretching bands originate from gradual changes in crystal structural unit cell parameters and

volumes due to the systematic Shannon effective nine-coordinated ionic radius decrease in the lanthanide contraction phenomenon.^{61–63}

Terahertz absorbing properties and first-principles calculations

Room-temperature (RT) THz absorption spectra for powders of **LnAg** were measured using THz-TDS to observe the low-frequency phonons. The THz absorption spectra in the measurement range of 0.5–1.5 THz reveal an intensive and sharp absorption signal between 1.0 and 1.2 THz with a maximum of around 1.1 THz for **LaAg**, **CeAg**, **PrAg**, and **NdAg** (Fig. 3a and $S6^\dagger$). Notably, this feature undergoes a gradual transition towards double peaks with a shoulder peak appearing at a higher frequency (*ca.* 1.15 THz) for samples **SmAg**, **EuAg**, **GdAg**, **TbAg**, and **DyAg**. The consistency and reproducibility of the peak position as well as splitting were further confirmed through multiple THz absorption measurements involving samples of varying thicknesses (Fig. $S7^\dagger$). As the amount of **LnAg** samples placed between sample holders increases, the absorbance increases simultaneously without altering the peak position. The multiple reflections between different interfaces barely impact the intense THz absorption signals (Fig. $S8^\dagger$). For precise determination of the peak position, the spectra were fitted with a single Lorentzian function (**LaAg**, **CeAg**, **PrAg**, and **NdAg**) and two Lorentzian functions (peak 1:



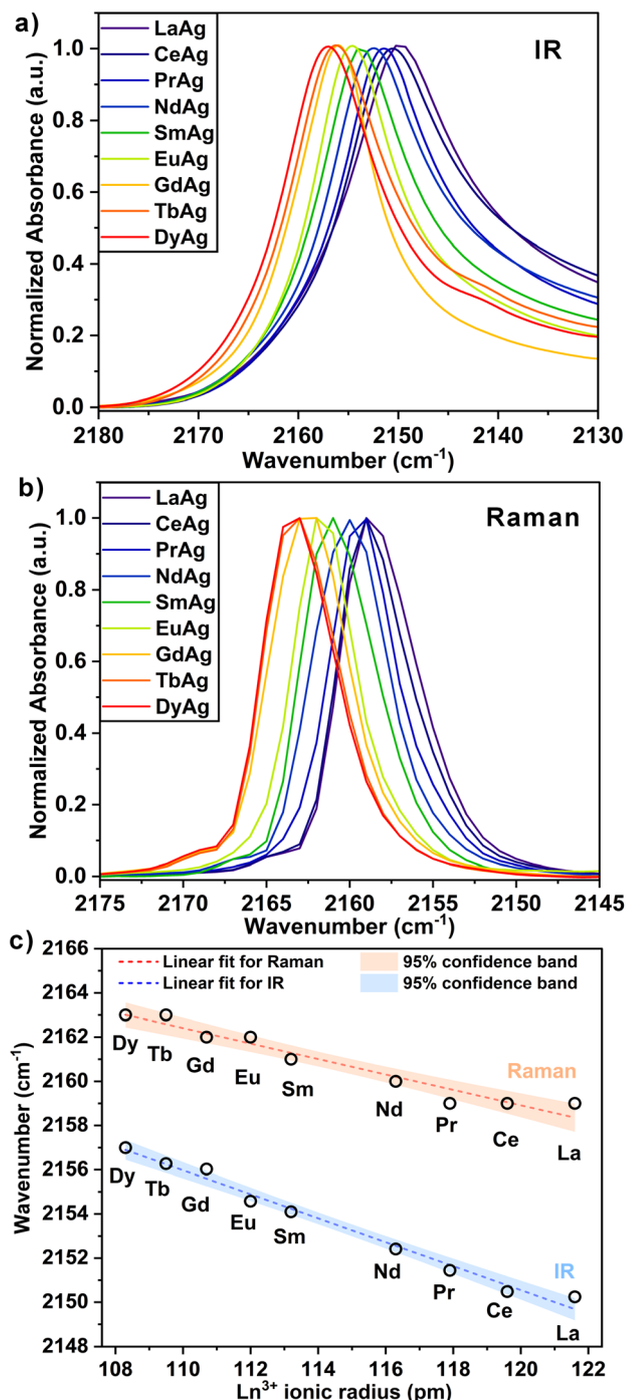


Fig. 2 Normalized IR absorption (a) and Raman scattering (b) spectra of LnAg in the CN⁻ stretching region. (c) Peak positions of the IR (blue) and Raman (orange) CN⁻ stretching band for different LnAg samples analyzed by linear fitting with 95% confidence bands.

higher frequency, peak 2: lower frequency) for SmAg, EuAg, GdAg, TbAg, and DyAg (Fig. 3c, d and Fig. S9, Table S4[†]). The THz spectral fitting unveils a gradual shift from the single peak at 1.090 THz for LaAg to peak 1 at 1.167 THz and peak 2 at 1.093 THz for DyAg. The approximately linear increase up to

74 GHz (0.074 THz) in frequency for both the single peak and peak 1 aligns coherently with the aforementioned IR spectra analysis (Fig. 3b). The low-temperature (liquid nitrogen, LN, $T = 77$ K) THz absorption spectra show a significant blueshift to a higher frequency for all LnAg samples (Fig. S10 and Table S4[†]). In particular, the most noticeable shift occurs for the PrAg sample with a change of 42 GHz, while LaAg, TbAg, and DyAg exhibit relatively modest responses of 14 GHz, 19 and 18 GHz (peaks 1 and 2), 19 and 24 GHz (peaks 1 and 2), respectively. Overall, the observed blue shifts, influenced by both lanthanide(III) ions and temperature-dependent factors, are attributable to the contraction in unit cell parameters and volume. Moreover, in comparison to room temperature, the full width at half maximum (FWHM) generally becomes narrower at low temperature for LnAg, aligning with the decrease in anharmonic thermal vibrations (Table S4[†]). The LnAg compounds with sharp absorbance together with narrow FWHM (ca. 100 GHz) show significant potential for providing efficient and selective shielding for THz radiations.

To further simulate these THz absorbances and gain deeper insights into their vibration origins, periodic DFT calculations of the phonon modes for LaAg and DyAg were performed starting from their experimentally determined crystal structures using the Vienna *ab initio* simulation package (VASP) and the PHONON code;^{66,67} see Experimental section for details. Previous reports have shown that first-principles calculations are effective tools to understand the properties and mechanisms of functional materials.^{68–70} In the case of the present compounds, the computed phonon modes show comparable phonon distribution and consist of IR active phonons for LaAg and DyAg defined as modes a and b, respectively, at 1.024 THz (a-5), 1.035 THz (a-6), 1.095 THz (b-3), and 1.125 THz (b-4) within the 1.0–1.2 THz range (Fig. 3e and Table S5[†]). The prominent infrared intensity observed in modes a-5 and b-3 can be assigned as transverse phonon modes with an IR active E_{1u} irreducible representation, while modes a-6 and b-4 in much weaker intensity belong to longitudinal phonon modes with a mixture of 9A_{2u} + 15E_{1u} and 7A_{2u} + 15E_{1u}, respectively. Considering the table of irreducible representations of point group D_{6h} (6/mmm) outlined in Table S6,[†] E_{1u} and A_{2u} correspond to the linear (x, y) and z functions, respectively. Notably, the higher frequency for DyAg phonon modes compared to LaAg phonon modes aligns with the blueshift in peak positions from LaAg to DyAg. Moreover, the substantial disparity in frequency (30 GHz) between phonon modes b-3 and b-4 for DyAg, in contrast to the 11 GHz difference for a-5 and a-6 in LaAg phonon modes, concurs with the experimentally observed peak splitting into a double peak for the latter lanthanides. The relatively reliable mimic of the experimental peak positions at low temperatures confirms the validity of the DFT THz calculation. Further analysis of the phonon density of states (phonon DOS) and phonon dispersion for LaAg and DyAg indicates similar distributions of partial DOS of atoms due to the isostructural nature of these compounds (Fig. 3f and S13, S14[†]). Particularly, in the low-frequency region, the largest contribution to the total phonon



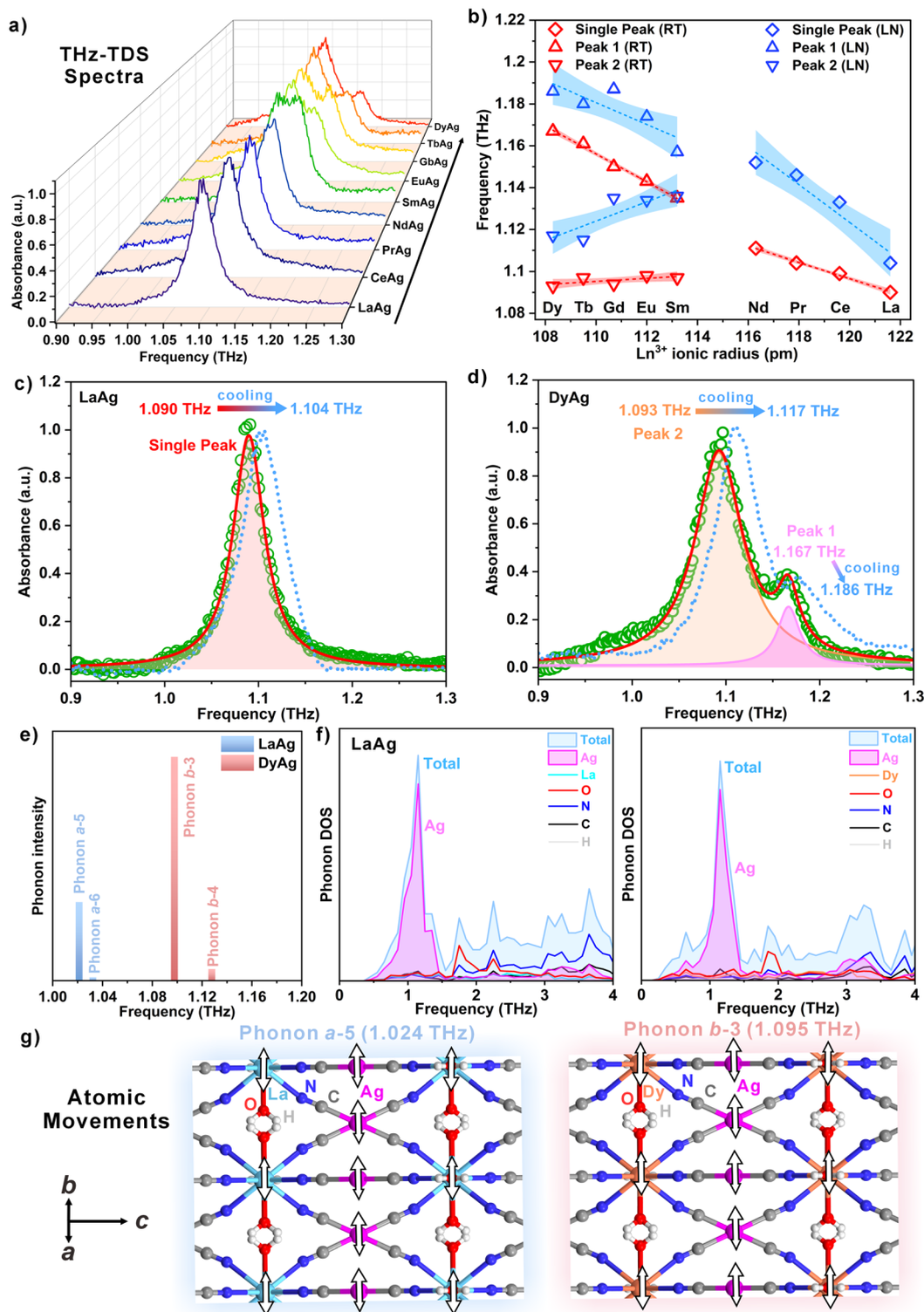


Fig. 3 (a) Room-temperature (RT) THz-TDS spectra of different LnAg samples after normalization. (b) Peak positions of THz-TDS spectra determined by fitted with Lorentzian function at RT and liquid nitrogen (LN) temperatures for LnAg samples. Dotted lines and colored areas represent linear fitting with 85% confidence bands. The normalized RT THz-TDS spectra of LaAg (c) and DyAg (d) fitted with Lorentzian functions. Red lines show the fitted curves, colored peaks show the components of each peak, and green dots indicate the experimental data. The blue dotted lines show the THz-TDS spectra at LN temperature. (e) Vertical bars showing the position and relative intensities of the IR active phonon modes for LaAg and b for DyAg at 1.024 THz (a-5), 1.035 THz (a-6), 1.095 THz (b-3), and 1.125 THz (b-4) in the indicated region. (f) Total phonon density of states (DOS) in light blue colored peaks of the LaAg (left) and DyAg (right) and partial phonon DOS of La, cyan line; Dy, orange line; Ag, pink colored peaks; O, red line; C, black line; N, blue line; H, grey line. (g) Atomic motion projections of the phonon modes for two main phonon modes a-5 and b-3. Color: La, cyan; Dy, orange; Ag, pink; C, dark gray; N, blue; O, red; H, light gray. The arrows indicate the vibration vectors.



DOS originates from Ag atoms, followed by O atoms. Consequently, the experimental THz absorbance observed at approximately 1.1 THz is primarily from the Ag vibrations within the networks. Based on the video of the atomic movements (Fig. 3g and Fig. S15, Movie S1†), the major phonon vibrations a-5 and b-3 for LaAg and DyAg exhibit similarity and can be interpreted as the opposite translational modes of Ag

atoms and H₂O molecules around Ln atoms in the *ab*-plane. This computational approach significantly enhances our comprehension of the intricate phonon dynamics governing the observed THz absorbances in LnAg.

Meanwhile, to investigate the THz wave absorbing capability, impedance matching analyses were conducted to simulate the return loss (RL) in decibels (dB) based on fitted single

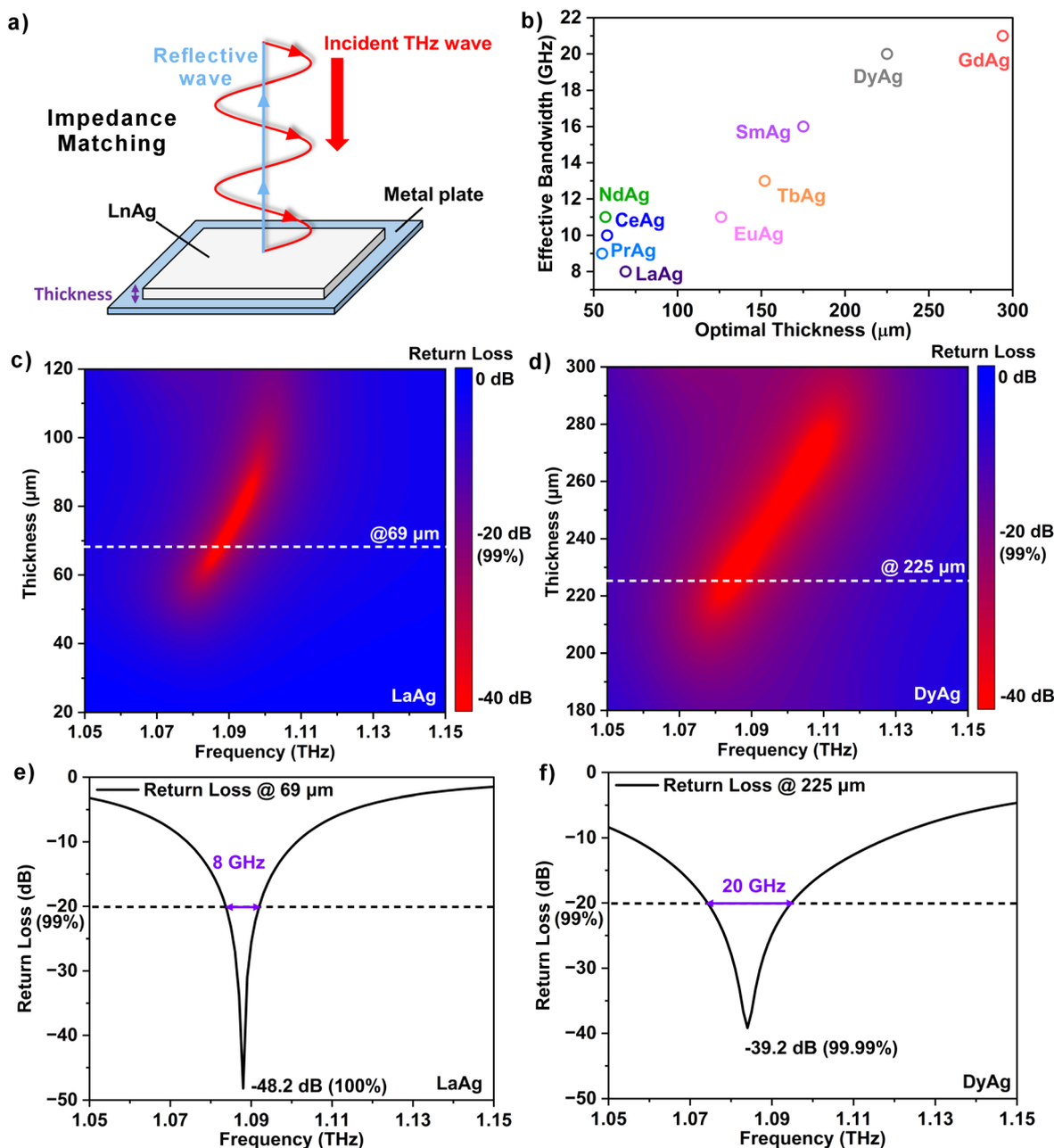


Fig. 4 Impedance matching analysis for the peak around 1.10 THz of LnAg. (a) Schematic illustration of THz absorber based on impedance matching. Irradiating an incident THz wave (red line) into LnAg absorbers on a metal plate minimizes the reflective wave (blue line) intensity in different thicknesses. (b) Comparison of THz wave absorbing performance for different LnAg samples. Colormaps depicting the RL intensity in dB as a function of frequency and thickness for LaAg (c) and DyAg (d) from impedance matching simulations. The white dash lines denote the optimal thickness with the largest RL intensity. The calculated RL spectra of the LaAg thin film (e) and DyAg thin film (f) from impedance matching simulations in the optimal thicknesses of 69 and 225 μm, respectively. The violet arrows mark the effective bandwidth (GHz) at -20 dB return loss.



Lorentzian function peak (**LaAg**, **CeAg**, **PrAg**, and **NdAg**) and the dominant peak 2 for **SmAg**, **EuAg**, **GdAg**, **TbAg**, and **DyAg** around 1.1 THz (Fig. 4 and S11, S12†),^{71–73} see Experimental section for details. As illustrated in Fig. 4a, the intensity of the reflected wave (blue line) diminishes across varying thicknesses when an incident THz wave (red line) is absorbed by **LnAg** absorbers on a metal plate. Remarkably, the minimum RL of -48.2 dB (100% absorption) and -39.2 dB (99.99% absorption) at 1.088 and 1.084 THz can be achieved with optimal 69 and 225 μm thicknesses for **LaAg** and **DyAg**, respectively. These optimal thicknesses correspond to the phase-matching condition, where the wave reflected on the surface of the absorber interferes with the wave reflected on the surface of the metal plate (Fig. 4c and d). In addition, based on the RL spectra at the optimal thicknesses, absorption intensities of -20 dB (99%) are satisfied over a wide range of 8 and 20 GHz for **LaAg** and **DyAg**, respectively (Fig. 4e and f); RL analyses for other **LnAg** samples are provided in the ESI (Fig. S11 and S12†). A comprehensive comparison of the THz wave absorbing performance for various **LnAg** samples is summarized in Fig. 4b. Generally, the results underscore that the **LnAg** compounds show an increase in effective bandwidth concerning thickness from **LaAg** to **DyAg**. In particular, **GdAg** possesses a superior effective bandwidth in 21 GHz, whereas the thinnest matching thickness (55 μm) can be achieved for **PrAg**. The simulations suggest that thin **LnAg** films possess a promising absorption performance at around 1.09 THz with a substantial return loss (<-20 dB) value across the maximum 21 GHz frequency range.

Compared to other state-of-the-art THz EM interference shielding materials in the literature, such as graphene foam, polymer composites, and two-dimensional transition metal carbides or nitrides materials, they usually reveal very broad THz shielding properties ranging from 0.2 to 2 THz.^{74–76} This work presents an easy preparation for a series of coordination complexes with a sharp and intense THz shielding performance exclusively around 1.09 THz with a narrow range. The tunable THz shielding effect from lanthanides allows the fine blocking of THz radiation in a specific and narrow range as a potential candidate for adjustable THz filters. Moreover, it should be noted that the wide optical transparency of most of **LnAg** was determined using solid-state UV-Vis-NR absorption spectroscopy (Fig. S4†), which offers the feasibility of optical window while shielding THz radiations simultaneously and presents potential applications for future THz devices.

Conclusions

In this work, a sequence of nine $[\text{Ln}^{\text{III}}(\text{H}_2\text{O})_3][\text{Ag}^{\text{I}}(\text{CN})_2]_3$ (**LnAg**, Ln = La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, and Dy) complexes were prepared and investigated their tunable vibrational spectra and temperature-dependent THz absorption properties systematically. It is noteworthy that all **LnAg** samples exhibit robust and adjustable absorptions with narrow FWHM, demonstrating significant responsiveness to variations in lanthanide(III)

ion or temperature in the range of 1.0 to 1.2 THz and displaying commendable reproducibility. Impedance matching calculations support the feasibility of preparing thin THz absorbing films with thicknesses less than 0.3 mm, yielding a significant return loss value in a broad frequency range.

In addition, advanced periodic DFT calculations based on the single-crystal structures of **LaAg** and **DyAg** reliably reproduce the experimental spectra within the computation approximation and highlight the immense potential to facilitate THz-absorbing materials by modifying the structural elements identified in the theoretical calculations. Further modulations may be realized by replacing the Ln(III) ions with other transition metals or by modifying the dicyanoargentate with other building blocks to meet the requirements of beyond 5G communication technologies. This comprehensive study opens up a novel and facile pathway for the design and tuning effects of THz waves, significantly contributing to our understanding of vibrations in the low-frequency domain.

Experimental

Materials

All starting materials: lanthanum(III) chloride heptahydrate, cerium(III) chloride heptahydrate, praseodymium(III) chloride heptahydrate, neodymium(III) chloride hexahydrate, samarium(III) chloride heptahydrate, europium(III) chloride hexahydrate, gadolinium(III) chloride hexahydrate, terbium(III) chloride hexahydrate, dysprosium(III) chloride hexahydrate, and potassium dicyanoargentate(I) were purchased from Sigma-Aldrich and Wako Pure Chemical Industries and utilized without further purification.

Caution! The cyanide compounds are toxic and should be handled with care.

General synthetic procedures for $[\text{Ln}^{\text{III}}(\text{H}_2\text{O})_3][\text{Ag}^{\text{I}}(\text{CN})_2]_3$ (**LnAg**, Ln = La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, and Dy) complexes

The reaction of a 0.3 mmol aqueous solution of $\text{LnCl}_3 \cdot 6\text{H}_2\text{O}$ (3 mL) with a 0.9 mmol solution of $\text{K}[\text{Ag}(\text{CN})_2]$ in an H_2O solution (3 mL) with vigorous stirring for 3 hours resulted in the formation of precipitation, from which the powder solids were collected by filtration under normal pressure and dried under the ambient conditions. The remaining solutions of **LaAg** and **DyAg** were evaporated slowly at room temperature inside a dark chamber to avoid photodecomposition and their crystals formed in a week. The crystalline and powder materials were used for further investigations. In addition, it should be noted that an analogous lanthanoid series of elements, specifically Y, Ho, Er, Tm, Yb, and Lu, was employed to synthesize similar compounds incorporating $[\text{Ag}(\text{CN})_2]^-$ ions based on identical synthetic procedures. However, the resultant samples predominantly comprised decomposed AgCN, associated with the smaller Ln(III) ion radius and the consequential collapse of the networks, which will not be considered in this work.

$[\text{La}^{\text{III}}(\text{H}_2\text{O})_3][\text{Ag}^{\text{I}}(\text{CN})_2]_3$ (**LaAg**). Yield: 89 mg (44% based on Ag). Anal. calcd for $\text{LaAg}_3\text{C}_6\text{H}_6\text{N}_6\text{O}_3$ (molar mass = 672.66 g



mol⁻¹): C, 10.71%; H, 0.90%; N, 12.49%. Found: C, 10.71%; H, 0.90%; N, 12.49%. FT-IR (KBr, cm⁻¹): 3513b, 1610s [$\nu(\text{O-H})$], 2150vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2159vs [$\nu(\text{C}\equiv\text{N})$].

[Ce^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**CeAg**). Yield: 93 mg (46% based on Ag). Anal. calcd for CeAg₃C₆H₆N₆O₃ (molar mass = 673.87 g mol⁻¹): C, 10.69%; H, 0.90%; N, 12.47%. Found: C, 10.79%; H, 0.98%; N, 12.44%. FT-IR (KBr, cm⁻¹): 3516b, 1610s [$\nu(\text{O-H})$], 2150vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2159vs [$\nu(\text{C}\equiv\text{N})$].

[Pr^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**PrAg**). Yield: 140 mg (69% based on Ag). Anal. calcd for PrAg₃C₆H₆N₆O₃ (molar mass = 674.66 g mol⁻¹): C, 10.68%; H, 0.90%; N, 12.46%. Found: C, 10.93%; H, 0.95%; N, 12.35%. FT-IR (KBr, cm⁻¹): 3521b, 1609s [$\nu(\text{O-H})$], 2151vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2159vs [$\nu(\text{C}\equiv\text{N})$].

[Nd^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**NdAg**). Yield: 121 mg (59% based on Ag). Anal. calcd for NdAg₃C₆H₆N₆O₃ (molar mass = 678.00 g mol⁻¹): C, 10.63%; H, 0.89%; N, 12.40%. Found: C, 10.70%; H, 0.91%; N, 12.33%. FT-IR (KBr, cm⁻¹): 3524b, 1609s [$\nu(\text{O-H})$], 2152vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2160vs [$\nu(\text{C}\equiv\text{N})$].

[Sm^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**SmAg**). Yield: 111 mg (54% based on Ag). Anal. calcd for SmAg₃C₆H₆N₆O₃ (molar mass = 684.12 g mol⁻¹): C, 10.53%; H, 0.88%; N, 12.28%. Found: C, 10.59%; H, 0.87%; N, 12.18%. FT-IR (KBr, cm⁻¹): 3528b, 1608s [$\nu(\text{O-H})$], 2154vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2161vs [$\nu(\text{C}\equiv\text{N})$].

[Eu^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**EuAg**). Yield: 110 mg (53% based on Ag). Anal. calcd for EuAg₃C₆H₆N₆O₃ (molar mass = 685.72 g mol⁻¹): C, 10.51%; H, 0.88%; N, 12.26%. Found: C, 10.58%; H, 0.89%; N, 12.18%. FT-IR (KBr, cm⁻¹): 3532b, 1608s [$\nu(\text{O-H})$], 2155vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2162vs [$\nu(\text{C}\equiv\text{N})$].

[Gd^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**GdAg**). Yield: 133 mg (64% based on Ag). Anal. calcd for GdAg₃C₆H₆N₆O₃ (molar mass = 691.00 g mol⁻¹): C, 10.43%; H, 0.88%; N, 12.16%. Found: C, 10.62%; H, 0.88%; N, 12.15%. FT-IR (KBr, cm⁻¹): 3535b, 1609s [$\nu(\text{O-H})$], 2156vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2162vs [$\nu(\text{C}\equiv\text{N})$].

[Tb^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**TbAg**). Yield: 106 mg (51% based on Ag). Anal. calcd for TbAg₃C₆H₆N₆O₃ (molar mass = 692.68 g mol⁻¹): C, 10.40%; H, 0.87%; N, 12.13%. Found: C, 10.51%; H, 0.84%; N, 12.11%. FT-IR (KBr, cm⁻¹): 3537b, 1609s [$\nu(\text{O-H})$], 2156vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2163vs [$\nu(\text{C}\equiv\text{N})$].

[Dy^{III}(H₂O)₃][Ag^I(CN)₂]₃ (**DyAg**). Yield: 38 mg (40% based on Ag). Anal. calcd for DyAg₃C₆H₆N₆O₃ (molar mass = 696.25 g mol⁻¹): C, 10.35%; H, 0.87%; N, 12.07%. Found: C, 10.47%; H, 0.83%; N, 12.10%. FT-IR (KBr, cm⁻¹): 3540b, 1609s [$\nu(\text{O-H})$], 2157vs [$\nu(\text{C}\equiv\text{N})$]. Raman (crystal, cm⁻¹): 2163vs [$\nu(\text{C}\equiv\text{N})$].

X-Ray crystallography

Pristine crystals of **LaAg** and **DyAg** submerged in Paratone (Hampton Research) oil, were mounted with a 100 μm Dual Thickness MicroMount™ (MiTeGen) loop and cooled down to 90 K in a cold nitrogen gas stream. X-ray diffraction data were collected with a Rigaku R-Axis RAPID diffractometer equipped with an imaging plate area detector using graphite monochromated Mo-K α radiation [$\lambda = 0.71075 \text{ \AA}$]. Data integration was performed by Rigaku RAPID AUTO software. Structures were solved by direct methods using SHELXS-97 incorporated in the Crystal Structure 4.0 crystallographic software package and

refined using an F^2 full-matrix least squares technique of SHELXL-2019/3 included in the OLEX-2 1.5 software package.^{77,78} Anisotropic refinement was performed for all non-hydrogen atoms, while hydrogen atoms were positioned with the help of the electron density map. Due to the presence of two heavier elements, La and Ag, and Dy and Ag for **LaAg** and **DyAg**, respectively, in the densely packed crystal structure, we observed strong diffuse scattering of X-rays with smaller density unassignable Q peaks. All attempts to solve this issue by introducing several disorder models did not improve refinement parameters. Therefore, residual density appeared due to the presence of heavier elements and defects in the crystal. Crystal data, data collection, and refinement parameters were gathered in ESI (Table S1†). CCDC numbers 2339311 and 2339312 contain the supplementary crystallographic data for **LaAg** and **DyAg**, respectively.† The structural models presented as figures were prepared with the use of the CCDC Mercury 3.10 visualization software.⁷⁹ Powder X-ray diffraction patterns at room temperature were collected for powdered samples placed on Si plates with a RIGAKU MiniFlex diffractometer equipped with monochromated Cu-K α radiation ($\lambda = 1.5406 \text{ \AA}$).

Physical characterizations

Elemental analyses (C, H, and N) were conducted with an Elementar Analysensysteme GmbH: vario MICRO cube. All infrared absorption spectra were collected with a JASCO FT/IR-4100 spectrometer for samples dispersed in the KBr pellet. JASCO NRS-7500 laser Raman spectrometer, equipped with a 532.12 nm laser (power = 0.8 mW), single monochromator, MPLFLN 20 \times /100 \times objective lens, BS 30/70, and 1–5% (OD 1.3) attenuator, was used to perform Raman spectroscopy in the range of 4000–100 cm⁻¹ (L 600/ B 500 nm grating, 10 \times 1000 μm slit, d -4000 μm aperture) on the single crystals of all samples in the reflectance mode. The terahertz (THz) wave absorption measurements at room temperature were conducted by compressing samples into pellets (diameter = 1.00 cm) of different thicknesses and masses between two white polyethylene plates using Advantest TAS7400TS THz-TDS in the transmittance mode. The thicknesses were determined five times for different pellet positions by the electronic outside micrometer with 0.001 mm resolution. For the THz-TDS measurements at liquid nitrogen temperature, the pellets of samples between two polyethylene plates were placed in a liquid-nitrogen-cooled cryostat (Oxford Instruments) connected to the digital temperature controller. Thermogravimetric analyses were performed on a Rigaku Thermo Plus TG8120 in the 25–400 $^{\circ}\text{C}$ range in dry air with a 2 $^{\circ}\text{C min}^{-1}$ heating rate and Al₂O₃ as reference material.

First-principles calculations

Periodic structure calculations were performed for **LaAg** and **DyAg** utilizing experimental crystal structures by using the Vienna *ab initio* Simulation Package (VASP).^{66,67} The 1 \times 1 \times 1 structures were fully optimized with convergence energy of 1.0 $\times 10^{-9}$ eV and the maximal residual forces on each atom less than 0.0001 eV \AA^{-1} . The optimized 1 \times 1 \times 1 structures of **LaAg**



and **DyAg** are used for further phonon calculations by the direct method implemented in the Phonon code with displacements of 1 pm.⁸⁰ The correlation energy was calculated using the generalized gradient approximation developed by Perdew, Burke, and Ernzerhof (GGA-PBE).⁸¹ A plane-wave basis set was used with a cut-off energy of 400 eV throughout the calculations. The k -mesh of $3 \times 3 \times 1$ for both **LaAg** and **DyAg** were used for the Brillouin zone samplings.

Impedance matching analysis

In the impedance matching analysis, a metal-backed THz wave absorber model composed of **LnAg** was designed.^{72,73} When a THz wave is vertically incident to a **LnAg** sheet applied on a reflective metal plate, interference occurs between the electromagnetic wave reflected on the surface of the sheet and the metal plate. The suitable thickness was simulated that exhibits maximum absorption by canceling out the reflected wave on the metal plate with that on the surface of the absorbers. Such a decrease in the reflection intensity is expressed by the return loss (RL) [dB], which is the amount of light reflected toward the source. It can be estimated through eqn (1) by using the normalized input impedance, Z_{in} .

$$RL [dB] = 20 \log_{10} \left| \frac{Z_{in} - 1}{Z_{in} + 1} \right| \quad (1)$$

$$Z_{in} = \sqrt{\frac{\mu_{new}}{\epsilon_{new}}} \tanh \left(j \frac{2\pi f d}{c} \sqrt{\epsilon_{new} \mu_{new}} \right) \quad (2)$$

where j , f , d , and c refer to the imaginary unit, frequency [Hz], sample thickness, and speed of light ($= 3.00 \times 10^8$ m s⁻¹), respectively. The permittivity $\epsilon_{new}(f)$ of the measured sample was obtained as follows, and the permeability $\mu_{new}(f)$ was set to 1.

As for obtaining the permittivity $\epsilon_{new}(f)$ in eqn (2), the observed THz wave absorption spectra for compounds **LnAg** were fitted with Lorentzian functions. The fitted single Lorentzian function peak (**LaAg**, **CeAg**, **PrAg**, and **NdAg**) and Lorentzian function peak 2 (lower frequency) for **SmAg**, **EuAg**, **GdAg**, **TbAg**, and **DyAg** around 1.1 THz were used for the impedance matching analyses. The frequency-dependent permittivity $\epsilon(f)$ of the compound was calculated by deciding ϵ_0 , ϵ_∞ , and calculated peak frequency f_p in eqn (3), which gave the minimum difference between the calculated absorption and the decomposed peak. The ϵ_0 and ϵ_∞ are the permittivities when the frequency is close to 0 and infinite, respectively.

$$\epsilon(f) = \epsilon_\infty + \frac{\epsilon_0 - \epsilon_\infty}{1 + 2j \left(\frac{\delta_p}{f_p} \right) \left(\frac{f}{f_p} \right) - \left(\frac{f}{f_p} \right)^2} \quad (3)$$

where δ_p is the attenuation of the calculated peak.

Calculated absorption S was obtained from eqn (4).

$$S [dB] = -20 \log_{10} \left(\frac{(1 - \Gamma(f)^2)T(f)}{1 - \Gamma(f)^2 T(f)^2} \right) \quad (4)$$

where $\Gamma(f)$ and $T(f)$ indicate the reflectance coefficient and transmittance coefficient obtained by the following eqn (5) and (6), respectively.

$$\Gamma(f) = \frac{\sqrt{\frac{\mu(f)}{\epsilon(f)}} - 1}{\sqrt{\frac{\mu(f)}{\epsilon(f)}} + 1} \quad (5)$$

$$T(f) = \exp \left(-j \frac{2\pi f d}{c} \sqrt{\epsilon(f) \mu(f)} \right) \quad (6)$$

In eqn (5), frequency-dependent permeability $\mu(f)$ was set to 1. The permittivity $\epsilon_{new}(f)$ of the measured sample was calculated by applying eqn (7). V_{air} and V_3 are the fill rate of air and **LnAg** in the measured pellet.

$$\epsilon_{new}(f) = V_{air} \epsilon_{air} + V_3 \epsilon(f) \quad (7)$$

Using the calculated permittivity $\epsilon_{new}(f)$ in eqn (1), the optimal sample thickness was obtained with the change of thickness and frequency, and the minimum value of return loss was around 1.10 THz.

Author contributions

G. L. – conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, validation, visualization, writing – original draft. O. S. – conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, supervision, validation, visualization. K. K. – formal analysis, funding acquisition, investigation, methodology, validation, visualization. Y. M. – data curation, formal analysis, funding acquisition, investigation, validation, visualization. L. W. – formal analysis, funding acquisition, investigation, validation, visualization. K. N. – funding acquisition, validation, visualization. M. Y. – formal analysis, validation, visualization. N. F. C. – formal analysis, funding acquisition, supervision, validation, visualization. S. O. – conceptualization, data curation, funding acquisition, methodology, resources, project administration, supervision, validation, visualization.

Conflicts of interest

There are no conflicts to declare.

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