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# Systematic investigation of the magneto-electronic structure and optical properties of new halide double perovskites Cs<sub>2</sub>NaMCl<sub>6</sub> (M = Mn, Co and Ni) by spin polarized calculations†

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A cohesive study using density functional theory simulations is performed to reveal and understand the structural stability, optoelectronic and magnetic properties of Cs<sub>2</sub>NaMCl<sub>6</sub> (M = Mn, Co and Ni) halide double perovskites. The exchange-correlation potential, which is the only unknown parameter in the state-of-the-art formulism is determined through the well-known generalized gradient approximation and integration of the mBJ potential to it. The structural optimization, mechanical stability criteria and tolerance factor confirmed the stability of the double perovskites in a cubic structure with  $Fm\overline{3}m$ symmetry. The elastic constants endorsed the mechanical stability and justify the brittle character of these double perovskites. The spin polarized electronic band profile and behaviour of the dielectric constant and absorption coefficient in the spin up and down channels revealed the presence of halfmetallic nature in these materials. Moreover, herein, we have discussed the origin of the half-metallic gap and magnetism. The unpaired electrons in the crystal field splitted d-orbitals of the M-sited constituents are responsible for the half-metallic and magnetic character. The total magnetic moment was determined to be  $4\mu_B$ ,  $4\mu_B$  and  $1\mu_B$  for the Mn-, Co- and Ni-based double perovskites, respectively, with main contributions solely coming from the transition metal atoms. The perfect spin polarization at the Fermi level suggests the application of double perovskites in spintronic technology.

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## Introduction

Spintronic technology has become the subject of growing interest since it allows the utilization of both spin degree of freedom and charge of electrons simultaneously.1-3 Therefore, spintronics is beneficial for developing devices with new functionality having multiple applications. It offers potential advantages over conventional electronic devices, such as marginal reduction in power consumption, increase in processing speed and nonvolatility data storage.4,5 Materials demonstrating high spin polarization (SP) character are suitable candidates for this technology. The ratio of the density of states (DOS) in the spin-up and down orientations at the Fermi level quantifies the magnitude of spin polarization,<sup>6</sup> SP =  $(DOS_{\uparrow} - DOS_{\downarrow})/(DOS_{\uparrow} + DOS_{\downarrow})$ . Diamagnetic and paramagnetic materials have symmetric DOS in the spin-up and down channels, and thus do not exhibit the spin polarization effect. However, ferromagnetic materials demonstrate asymmetric DOS, and thus their SP is always greater than zero but less than unity. In the case of a half-metallic materials:

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band structure in one spin channel is metallic showing the presence of states at the Fermi level, whereas exhibit semiconducting/insulting nature in the other spin channel. Therefore, these materials demonstrate 100% spin polarization at the Fermi level and are thought to be suitable candidates for spintronic technology.7-9 Pioneer work in field of half-metallicity was carried out by de Groot et al. in 1983.7 Since then, numerous compounds from different families have been reported to demonstrate half-metallic character. Among the variously reported half-metallic materials, the growing demand is for those materials that demonstrate high magnetic moment and high Curie temperature.

Perovskites exhibit a stoichiometric formula of ABX3 with A as s/p element, B is mostly a transition/inner-transition element or even can belong to the s/p block, while X can be either oxygen (oxide perovskites) or halogen member (halide perovskites). 10,11 The perovskite family is probably the most studied family because they can be synthesized and exhibit diverse physical properties, such as radioactive waste encapsulation, high-temperature superconductivity, and multiferroicity, which make these ceramics an imperative tool for various fascinating technologies, including solid-oxygen fuel cells, spintronics, and shape memory devices. 12-14 Furthermore, the flexibility of the structure of perovskites allows easy

tailoring/modification of their stoichiometry to optimize their electronic, magnetic and thermoelectric properties. Over the past few decades, there has been particular interest in doubling the constituents of perovskites, while replacing exactly half of the B-site cations by another cation (B'), forming the so-called double-perovskite A<sub>2</sub>BB/X<sub>6</sub>. <sup>15</sup> Due to the interplay between the B- and B'-sited cations, double perovskites show half-metallicity and other feasible properties. Oxide double perovskites have been reported to depict fascinating properties. Sr<sub>2</sub>FeReO<sub>6</sub>, Sr<sub>2</sub>FeMoO<sub>6</sub>, Sr<sub>2</sub>CrWO<sub>6</sub> and their related double perovskites are half-metals with a fairly high magnetic transition temperature, making them suitable candidates for future spintronics applications. 16,17 Ba<sub>2</sub>FeMoO<sub>6</sub> has a figure of merit (ZT) nearly equal to unity, and thus is suitable for thermoelectric technology. 18 On the other hand, halide perovskites (X belonging to the halogen family) have been used to produce optoelectronic devices with impressive quality.19 These materials have revolutionized solar cell technology by achieving an efficiency of about 22%.19,20 However, recently Cai et al. investigated some lead-free Cs<sub>2</sub>GeB'X<sub>6</sub>-type halide perovskites and reported that these materials exhibit ferromagnetic character with a high Curie temperature.21 Moreover, E. Haque and M.A. Hossain investigated Cs<sub>2</sub>BiAgX<sub>6</sub> (X = Cl, Br) materials and found the presence of ultralow thermal conductivity in these materials, giving rise to  $ZT \approx 0.8^{22}$  Therefore, these materials are expected to show a good thermoelectric response.

In the current study, we present a detailed report of new possible half-metallic halide double perovskites. The motive of the present study is to explore the structural stability and understand the origin of half-metallicity in these materials. The electron filling in the d-states of transition atoms plays a vital role in opening the semi-conducting gap in one of the spin directions. Numerous alkali metal halide Cs2NaB'Cl6-type double perovskites have been reported since the 1970s. 23-26 The focus of the investigations was mostly to understand the structural stability of those compounds. Among them, most of the materials such as Cs<sub>2</sub>NaScCl<sub>6</sub>, Cs<sub>2</sub>NaTiCl<sub>6</sub>, Cs<sub>2</sub>NaVCl<sub>6</sub>, Cs<sub>2</sub>NaFeCl<sub>6</sub>, Cs<sub>2</sub>NaAmCl<sub>6</sub>, and Cs2NaBkCl6 were found to be crystalline in cubic structures following symmetric operations of the  $Fm\bar{3}m$  space group. Within the crystalline structure, cesium is enclosed by a cage of 12-Cl atoms, while an Na/B' atom lies within the octahedron environment of Cl. However, to date, their electronic band profile,

magnetic character and transport properties still remain unexplored. Thus, based on DFT calculations, herein, we present the structural, mechanical, electronic, optical and magnetic properties of  $\mathrm{Cs_2NaMCl_6}$  (M = Mn, Co and Ni) double perovskites.

#### **Computational details**

Density functional theory (DFT) and post-DFT simulations have proven to be one of the most accurate methods for the computation of the structural magneto-electronic structure of crystalline materials.27,28 The ground state structural parameters of the titled double perovskites are predicted by optimization and relaxation of their unit cell structure. The optimized structure parameters are used to predict the electronic and optical profile of the double perovskites. All the calculations are performed with the help of first principle methods, wherein we have solved the Kohn-Sham equation through an iterative scheme with the help of the Wein2k code and simulation packages integrated to it. The presence of delectrons of Mn/Co/Ni may increase the exchange correlation effects in the Cs2NaMCl6 double perovskites. Thus, the generalized gradient approximation (GGA) without and with the modified Becke-Johnson potential (GGA + mBJ) is used. 29,30 Muffin tin spheres are framed around atomic constituents. The LAPW basis set is incorporated for the wave expansion, which ensured that the waves within the sphere are atomiclike and outside it has plane wave character. The parameters  $l_{\text{max}} = 10$  and  $R_{\text{MT}}K_{\text{max}} = 7$  are used for the atomic and plane wave cutoff, respectively. A reduced Brillouin zone (BZ) scheme that could simulate the crystal under periodic boundary conditions is used. For the integration over the BZ, a compact mesh of 1000 k-points is incorporated. The iterations stopped self consistently when the convergence conditions 0.0001e for charge and 0.0001 Ry for energy are satisfied. The elastic constants and optical properties, which can help in understanding the mechanical stability and exploration of electronic structure are determined by incorporating the elastic and optic package implemented with the Wein2k simulation code.

#### Structural stability

Structural stability is crucial for any material, which affects almost all its physical properties. The titled double perovskites have not been synthesized experimentally to date; however,

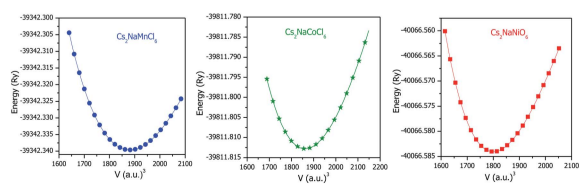


Fig. 1 Optimization curves of Cs<sub>2</sub>NaM(Mn, Co, Ni)Cl<sub>6</sub> double perovskites by spin polarized calculations.

 $Cs_2NaB'Cl_6$  (B' = Sc, Ti, V and Fe) has been prepared, exhibiting a cubic structure having symmetry of the  $Fm\bar{3}m$  space group. <sup>25,26</sup> By employing simulation, the stability and ground state structure of any material can be explored by minimizing its crystal energy with respect to volume. Sequentially, to establish the ground state structure of the Cs<sub>2</sub>NaMCl<sub>6</sub> (M = Mn, Co and Ni) double perovskites we optimized their structures with spin polarized calculations. The optimization curves are represented in Fig. 1, which disclose the stability of these double perovskites. The minimum in the E-V curve corresponds to the ground state energy. The equilibrium lattice parameters are evaluated at the optimized volume corresponding to the minimum energy and are tabulated in Table 1. On comparing the unit cell volume, it is evident that the lattice constant decreases in the order of Mn > Co > Ni-based double perovskites. This is because atomic size decreases across the period. The unit cell structures are shown in Fig. S1 (ESI),† where it can be seen the Na/M-atoms are within octahedra of Cl-atoms, while Cs is coordinated by 12 Cl-atoms.

computing the tolerance factor  $(\tau)$  as:<sup>31</sup>  $\tau = \left(\frac{0.71\langle \text{Cs} - \text{Cl} \rangle}{\langle \text{M}' - \text{Cl} \rangle_{\text{avg}}}\right)$ , where  $\langle \text{Cs-Cl} \rangle$  represents the inter-atomic distance between Cs and Cl atoms  $\langle \text{M}' - \text{Cl} \rangle_{\text{avg}}$  is the average of the  $\langle \text{Na-Cl} \rangle$  and  $\langle \text{M-Cl} \rangle$  bond distances. The condition  $\tau$  equals unity corresponds to an ideal double perovskite structure exhibiting the symmetry of the  $Fm\bar{3}m$  space group. The ideal structure is free from any internal stress and the Na-Cl-M bond angle is 180°. However,

the cubic phase is stable over the range of 0.9-1. When the

The stability of the cubic structure is further confirmed by

tolerance factor is smaller than the lower limit, the Na-Cl and M-Cl bonds remain compressed, while the Cs-Cl bonds are under tension. The structure relaxes and overcomes stress via the cooperative rotation of the octahedrons, which lowers the symmetry of the double perovskite. These rotations are meant to bend the Na–Cl–M bond angles from  $180^{\circ}$  to  $180^{\circ}$ – $\theta$  and the stability of the cubic phase decreases compared to lower symmetric phases. When  $\tau > 1$ , the Na–Cl and M–Cl bonds come under tension, whereas the Cs-Cl bonds are under compression, and under this condition the Na-Cl-M bond angle remains 180°; however, the structure overcomes the tensile stress by allowing hexagonal stacking. The bond length and bond angle of the optimized structure are reported in Table 1. The evaluated value of the tolerance factor from the bond lengths reveals the stability of double perovskites in the cubic structure.

#### Mechanical stability and properties

The mechanical stability and strength of materials can be explored by determining their elastic constants. The sufficient number of elastic constants necessary to survey the mechanical behaviour depends on the structural symmetry exhibited by a system. Since the double perovskites under investigation stabilize in cubic symmetry, therefore only three elastic constants,  $C_{11}$ ,  $C_{12}$  and  $C_{44}$ , are required to analyse their mechanical behaviour. Born formulated the necessary condition for the mechanical stability, which are given by  $C_{11} > 0$ ;  $C_{11} - C_{12} > 0$ ;  $C_{44} > 0$ ;  $C_{11} + 2C_{12} > 0$ . <sup>32,33</sup> The determined elastic constants in Table 2 follow Born's stability criteria, thereby

**Table 1** Ground state parameters of volume V (a.u.)<sup>3</sup>, crystal energy,  $E_0$  (Ry), cohesive energy  $E_{coh}$  (Ry), bond Length (a.u.), bond angle and tolerance factor ( $\tau$ )

				Bond length		Bond angle			
Double perovskite	V	$E_0$	$E_{\mathrm{coh}}$	Cs-Cl	Na-Cl	M-Cl	Cl-Cs-Cl	Na-Cl-M	τ
Cs <sub>2</sub> NaMnCl <sub>6</sub>	1880.86	-39342.3396	2.05	6.93	4.89	4.89	90°	$180^{\circ}$	0.99
Cs <sub>2</sub> NaCoCl <sub>6</sub> Cs <sub>2</sub> NaNiCl <sub>6</sub>	1861.74 1800.25	-39811.8128 $-40066.5840$	2.01 2.09	6.86 6.88	4.85 4.86	4.85 4.86	90° 90°	$180^{\circ}$ $180^{\circ}$	0.99 0.99

Table 2 Calculated elastic parameters at 0 GPa and 0 K

Parameter (GPa)	$\mathrm{Cs_2NaMnCl_6}$	$\mathrm{Cs_2NaCoCl_6}$	Cs₂NaNiCl <sub>6</sub>
$C_{11}$	49.22	39.78	36.80
$C_{12}$	18.11	16.24	15.52
$C_{44}$	17.23	15.51	14.98
$B = B_{\rm V} = B_{\rm R} = (C_{11} + 2C_{12})/3$	28.48	24.08	22.25
$G_{\rm V} = (C_{11} - C_{12} + 3C_{44})/5$	16.56	14.41	13.65
$G_{\rm R} = 5C_{44}(C_{11} - C_{12})/\{4C_{44} + 3(C_{11} - C_{12})\}$	16.51	14.28	13.44
$G = (G_{\rm V} + G_{\rm R})/2$	16.54	14.34	13.54
Y = 9BG/(3B+G)	41.57	36.07	34.04
B/G	1.72	1.72	1.71
$\sigma = (3B - Y)/6B$	0.25	0.25	0.25
$A = 4C_{44}/(C_{11} - C_{12})$	1.10	1.21	1.28
$H = 0.92(B/G)^{1.3137}G^{0.708}$	20.12	18.56	17.44

authenticating the mechanical stability of the double perovskites. Furthermore, the elastic constants can provide insight to mechanical behaviour through the computation of the elastic moduli and other parameters. The Reuss and Voigt values of the bulk (B) and shear (G) moduli represent the extreme ends of the bulk and shear moduli.<sup>34</sup> Hill's scheme is a practical way for estimating these moduli by averaging of the extreme values, *i.e.*,  $B = (B_V + B_R)/2$  and  $G = (G_V + G_R)/2$ , where  $B_V$  and  $G_V$  are the Voigt bulk and shear moduli, respectively, which can be calculated directly from the elastic constants using the relation  $B_V = (C_{11} + C_{12})/3$ ;  $G_V = (C_{11} - C_{12} + 3G_{44})/5$ .  $B_R$  and  $G_R$  are the Reuss values of the bulk and shear moduli, which can be obtained using the following relations:  $B_V = \frac{C_{11} + C_{12}}{3}$  and  $G_R = \frac{SC_{44}(C_{11} - C_{12})}{4C_{44} + 3(C_{11} - C_{12})}$ , respectively. Directly from the B and G values, one can deduce the Young's modulus

(Y) and Poisson's ratio ( $\sigma$ ) using the relations Y = 9BG/(3B + G)and  $\sigma = (3B - Y)/6B$ , respectively. The calculated values are listed in Table 2. Generally, the elastic modulus is the resistance offered by any material towards deformation, the bulk modulus is for volumetric deformation and the shear modulus for shape deformation, whereas Y characterizes the resistance to uniaxial deformation and can be used to estimate the stiffness of a material. The moduli decreased in the order of Y > B > G, signifying that these double perovskites allow shape deformation more easily compared to volumetric deformation. The mechanical behaviour of brittleness or ductility can be directly described from the elastic parameters. The different empirical parameters that are often used to signify mechanical behaviour are the Pugh ratio (B/G) and Poisson's ratio.<sup>3,34</sup> The index values separating the extreme behaviours are 1.75 and 0.26 for B/G and  $\sigma$ , respectively. Values above these decisive numerals for any

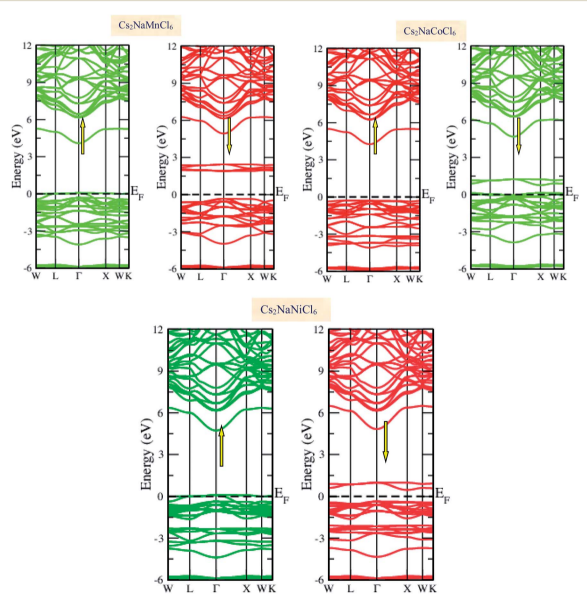


Fig. 2 Spin polarized band structures of  $Cs_2NaMCl_6$  double perovskites *via* GGA approximation. Red lines are used for semiconducting channels and green for metallic channels.

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material demonstrate the ductile character, otherwise brittle in nature. Accordingly, the calculated values reveal the brittle nature of the titled double perovskites. Additionally, the Zener anisotropic A factor is determined using the elastic constants and is given in Table 2. It used to characterize the deviation from isotropic behaviour. The calculated values suggest the presence of anisotropy since the magnitude of A deviates from unity. We also computed the micro-hardness (H) to examine the mechanical strength of the double perovskites. The hardness of a material can be directly linked to its bulk or shear modulus. The calculated values of H given in Table 2 suggest that all three Cs-based double perovskites have good mechanical strength.

#### Electronic and magnetic properties

Materials with smart electronic and magnetic profiles are enormously demanded in various scientific fields. These materials are not merely interesting for increasing fundamental knowledge, but also have implications for important industrial applications. Therefore, understanding the electronic and magnetic nature of materials is essential. Herein, we plotted the variation in the density of states with energy and also the band structures (E-k diagrams) with a reduced Brillouin zone scheme to outline the electronic character of the materials. The energy state distribution is greatly affected by the exchange and correlation potential. The presence of d-state electrons may make the GGA inefficient for these double perovskites. Thus, we also incorporated the mBJ potential to define a more precise band structure. The band structure of the double perovskites via GGA is demonstrated in Fig. 2 and the magnified view of the band structures is presented in Fig. S2 (ESI).† The behaviour of the E-k dispersion curves in the vicinity of the Fermi level designates the electronic properties of any material. The overlap of the partially filled energy bands with the Fermi level gives rise to metallic character. However, completely filled bands can be found beneath the Fermi level are responsible for opening the semiconducting band gap. From the band structure, it can be

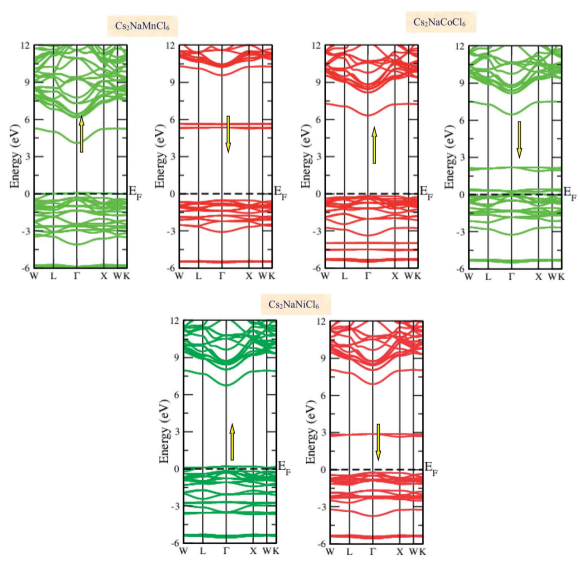


Fig. 3 GGA + mBJ band profiles of Cs<sub>2</sub>NaMCl<sub>6</sub> double perovskites

**Table 3** Observed band gap ( $E_{\rm g}$  in eV) in semi-conducting channel and total magnetic moment ( $M_{\rm T}$  in  $\mu_{\rm B}$ ) of Cs<sub>2</sub>NaMCl<sub>6</sub> double perovskites through the GGA and GGA + mBJ approximations

	Cs <sub>2</sub> NaMnCl <sub>6</sub>		Cs <sub>2</sub> NaCoCl <sub>6</sub>		Cs <sub>2</sub> NaNiCl <sub>6</sub>	
Approximation	$E_{ m g}$	$M_{ m T}$	$E_{ m g}$	$M_{ m T}$	$E_{ m g}$	$M_{\mathrm{T}}$
GGA GGA + mBJ	2.06 5.03	4.0 4.0	4.25 6.07	4.0 4.0	0.96 2.9	1.0 1.0

observed that Cs2NaMnCl6 and Cs2NaNiCl6 show metallic character in the spin-up channel and semiconducting nature in the opposite spin direction. However, Cs2NaCoCl6 demonstrates a semiconducting band structure in the spin-up direction and metallic character in the spin-down direction. The closeness of the Fermi level to the valance band in the semiconducting channel indicates the presence of p-type semiconducting behaviour. The existence of metallic character in one spin channel and semiconducting in the other induces halfmetallicity. Moreover, pseudo-gaps can be observed from the band profile of the double perovskites. In the metallic channel of Cs2NaMnCl6 and Cs2NaNiCl6, a forbidden gap originates from 0.17 eV to 4.2 eV and 0.15 eV to 5.2 eV, respectively. In contrast, in the spin-down channel of the Co-based halide perovskite, the forbidden gaps are in the region of 0.25 eV to 0.8 eV and 1.12 eV to 5.05 eV. The half-metallic character is maintained even when we incorporated the mBJ potential to GGA, as shown in Fig. 3. However, the range of the semiconducting gap is affected, as presented in Table 3.

The total DOS plots shown in Fig. 4 reveal the comparative effect of the mBJ potential on the electronic properties of the  $Cs_2NaMCl_6$  double perovskites. The electron states are shifted away from the Fermi level, thereby widening of band gap can be observed in the semiconducting channel. To further illustrate the band structure and clarify the energy states associated with the valance and the conduction bands, we plotted the partial density of states. The Cl-atom contributes the most towards the valance band formation, as shown in Fig. 5. In contrast, the belonging of the Cs and Na states are mostly away from the

valance band, and thereby do not play any significant role in shaping the electronic structure. The d-states of the transition atom play a peculiar role, and thereby dominate the electronic properties. On comparing the pDOS of the three double perovskites, it is evident that the d-states of the transition atoms are placed differently across the Fermi level are responsible for their half-metallic character. The p-d hybridization occurs among the dt<sub>2g</sub> states of Mn and Cl in the valance band of the spin-up channel. The Mn-deg states at the Fermi level give rise to the metallic character in the spin-up channel of the Mn-based double perovskite. In contrast, in the spin-down channel, all the d-states of Mn are above the Fermi level form the conduction band minima. In the Co-based double perovskite, both the dt<sub>20</sub> and deg states of the spin-up channel below the Fermi lever take part in p-d hybridization. On the other hand, the dt<sub>2g</sub> states are at the Fermi level in the spin-down channel and the deg states are empty lie above the Fermi level. However, in the Ni-based double perovskite, the dt<sub>2g</sub> states are completely filled for both spin channels and lie in the valance band. The deg states in the spin-up channel at the Fermi level give rise to metallic character, while in the spin-down state they form the conduction band minima. The p-states of the Cl-ligands are completely filled, and mainly contribute to the formation of the valance band. The valance states of Cs-s and Na-s are empty, and thus are located at higher energies above the Fermi level. The inner valance p-states of Cs give rise to the DOS peak at around -5 eV for both spin directions in these materials. Moreover, it can be analysed from the DOS plots that Cs, Na and Cl have equal amounts of electrons in the up- and down-states, having almost symmetric DOS across the reference level. The d-states of the transition atom are asymmetrically distributed, and therefore give rise to the magnetic character in the double perovskites.

## Origin of half-metallicity and ferromagnetic character

As mentioned in the structural properties, the M-site atoms are located in the octahedral environment of the Cl-atoms. The crystal field due to M–Cl Coulomb interactions decrease the degeneracy of the Mn/Co/Ni d(d $_{xy}$ , d $_{yz}$ , d $_{zx}$ , d $_{x^2-y^2}$ , d $_{z^2}$ ) orbitals, which are splitted into dt $_{2g}$  (d $_{xy}$ , d $_{yz}$ , d $_{zx}$ ) and de $_{g}$  (d $_{x^2-y^2}$ , d $_{z^2}$ ) sets. The dt $_{2g}$  state has an intake capacity of 6 electrons (3  $\uparrow$  + 3  $\downarrow$ ),

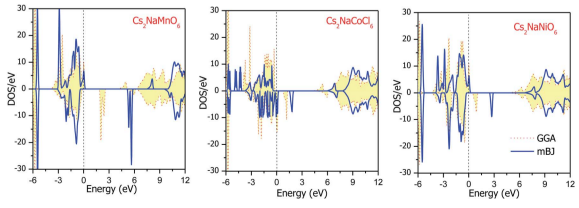
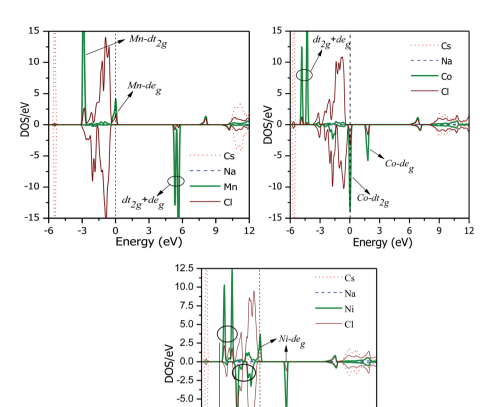


Fig. 4 Energy state distribution over the energy range of -6 to 12 eV by GGA and mBJ approximations.

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Fig. 5 PDOS of  $Cs_2NaMCl_6$  double perovskites in the energy range of -6 to 12 eV.

-7.5 -10.0

while the deg state can accommodate a maximum of 4 electron  $(2\uparrow + 2\downarrow)$  following Hund's rules. After half filling  $dt_{2g}$  orbitals, the electrons can choose to either form a high-spin state, i.e., fill higher energy deg before pairing occurs in the dt2g orbitals or to form a low-spin state i.e.,  $de_{\alpha}$  states are filled only after the  $dt_{2\alpha}$ states are completely filled. The choice of forming a high-spin state or low-spin state depends on the crystal field splitting energy (CFS) and pairing energies. The high-spin state is formed only if the splitting energy is less than the pairing energy. The CFS is affected by the number of electrons and size of the central atom.35 The type of ligand affects the splitting, strong field ligands can do large splitting. However, in the presence of weak field ligands such as Cl, the CFS is small and therefore the formation of the high-spin state depends on the oxidation number, electron number and size of the transition anion. The expected oxidation states in these double perovskites are Cs<sub>2</sub><sup>1+</sup>Na<sup>1+</sup>M<sup>3+</sup>Cl<sub>6</sub><sup>1-</sup>. Mn<sup>3+</sup>, Co<sup>3+</sup> and Ni<sup>3+</sup> have 4, 6 and 7 valance electrons, respectively, in their d states. The electron occupancy in the Mn<sup>3+</sup> d state is  $3t_{2g}(\uparrow)$ ,  $1e_g(\uparrow)$ ,  $0t_{2g}(\downarrow)$ , and  $0e_g(\downarrow)$ , forming a high-spin state with S = 2. The d states in the spin-down state are empty and form the conduction band minima. The  $de_{\mathrm{g}}$ state in the spin-up channel is partially filled and resides at the Fermi level, as is evident in Fig. 5. According to the DOS plots of the Co-based double perovskite, it is obvious that the d states in

the spin-up channel are completely occupied. The dt<sub>2g</sub> states in the spin-down channel are at the Fermi level. The electron filling in the valance states of the  $\text{Co}^{3+}$  states therefore is  $3t_{2g}(\uparrow)$ ,  $2e_g(\uparrow)$ ,  $1t_{2g}(\downarrow)$ , and  $0e_g(\downarrow)$ , thereby leaving four electrons unpaired with S = 2. The occupancy clearly confirms that the dorbitals in the spin-up channel that are completely filled form the valance band. While, the  $dt_{2g}$  states in the anti-parallel spin channel that are partially filled give the metallic character in the respective spin channel. Upon going from Mn to Ni, the number of electrons at the central atom increases, while size decreases.  $Ni^{3+}$  has the electronic configuration of  $3t_{2g}(\uparrow)$ ,  $3t_{2g}(\downarrow)$ ,  $1e_g(\uparrow)$ , and  $0e_g(\downarrow)$ , leaving one unpaired electron, and thus S = 1/2. Thereby, the spin state changes from high to low when Ni is the central atom. Each unpaired electron gives a magnetic moment of  $\approx 1\mu_{\rm B}$ , and thus the total magnetic moment becomes  $4\mu_B$ ,  $4\mu_B$  and  $1\mu_B$  for the Mn-, Co- and Ni-based double perovskites, respectively. The integral magnetic moment also signifies the presence of half-metallicity.

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#### Nature of bonding from charge density

The characteristic of bonding can be determined from the charge density and its spatial distribution. The ionic bond is characterised by a charge-free inter-atomic distance, whereas metallic and covalent bonds have charge sharing among constituents.<sup>31</sup>

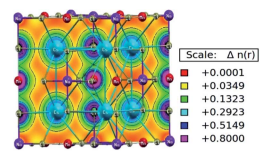


Fig. 6 Spatial charge distribution of  $Cs_2NaMnCl_6$  double perovskites along the body diagonal.

However, in the covalent bond, charge sharing is directional, whereas the metallic bond has a uniform charge distribution throughout. The spatial charge configuration along the body diagonal plane for the Mn-based double perovskite is depicted in Fig. 6. The maximum charge resides at the atomic positions. The Cs and Na atoms have a completely spherical charge distribution and there is no overlapping of charge contours with the neighbouring Cl atom. This signifies that the Cs and Na atoms have ionic bonding character with the Cl atom. Conversely, Mn displays covalent bonding character with the Cl atom since the charge distribution deviates from prefect spherical to deformed (dumb-bell type). The overlapping of the contours signifies the hybridization among the M–Cl orbitals. The p–d hybridization is also confirmed by the PDOS distribution.

#### **Optical properties**

The optical parameters of the Cs<sub>2</sub>NaMCl<sub>6</sub> double perovskites are calculated in order to analyze their possible optoelectronic applications and to gain deeper insight into their electronic

band profile. Since the band structures and DOS plots shown in Fig. 2 to 5 reveal metallic character in one spin channel and semiconducting in the other channel, it is obvious that the optical response will be different in these channels. Thus, we have determined the spin polarized optical behaviour of the Cs<sub>2</sub>NaMCl<sub>6</sub> double perovskites. The complex dielectric function  $\varepsilon(\omega)$  is often used to completely portray the response of a material to disturbances due to electromagnetic radiation.  $\varepsilon(\omega)$  has two components,  $\varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega)$ , where the real part  $\varepsilon_1(\omega)$  defines the scattering power of a material. 36-38 Meanwhile, the imaginary part has a direct relation with the band structure and reflects the absorptive behaviour, which is quite helpful in material selection for solar cell technology. The dielectric constant components are interrelated through the Kramer-Kronig relationships.<sup>37</sup> The variation of  $\varepsilon_1(\omega)$  and  $\varepsilon_2(\omega)$  with photon energy is shown in Fig. 7(A and B), where it is clear that in the spin channels of Mn-up, Co-dn and Ni-up, the high value becomes negative between 0-1.2 eV for the Mn- and Ni-based double perovskites. Thereafter, they do not exhibit sharp variations with energy. In the negative range of  $\varepsilon_1(\omega)$ , incident electromagnetic radiation (photon) is completely attenuated by the double perovskites, revealing their perfect metallic nature. The points at  $\varepsilon_1(\omega)$  drop below unity and have a positive slope, which are known as plasmon excitation points. However, in the spin channels of Mn-dn, Co-up and Nidn,  $\varepsilon_1(\omega)$  increases from a static value limit (zero frequency). The static values increase in the order of 1.84, 2.17, and 3.01 for Co-up, Mn-dn, and Ni-dn, respectively. The variation in the static value from one spin channel to another and by changing magnetic ions is expected due to the inverse relation of  $\varepsilon_1(0)$ 

with the band  $\varepsilon_1(0)=1+\left(\frac{\hbar\omega}{E_{\rm g}}\right)$ . The imaginary part of the

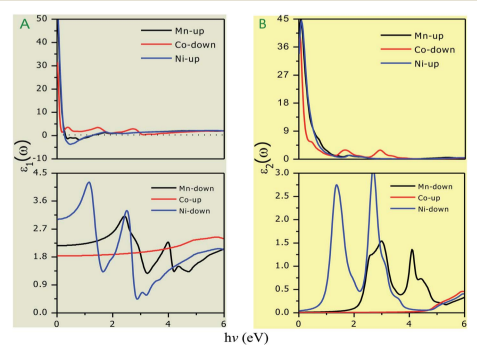


Fig. 7 Variation in dielectric constant with photon energy. (A) Real part ( $\epsilon_1$ ) with gray background and (B) imaginary part ( $\epsilon_1$ ) with yellowish background.

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1200 A B Mn-up Mn-up Co-dowr Co-down 900 Ni-up Ni-up 20 600 10 300  $\sigma(\Omega^{-1} cm^{-1})$  $\alpha (10^4 \, \text{Cm}^{-1})$ 30 1200 Mn-down Mn-dowr Co-up Ni-down 900 Ni-down 20 600

Fig. 8 (A) Variation in absorption coefficient (α) with photon energy with grey background and (B) optical conductivity with photon energy.

 $h_{V}$  (eV)

300

dielectric constant  $\varepsilon_2(\omega)$  is very important in signifying the efficiency of optoelectronic devices, representing the photons that can be absorbed by perovskites. The optical edge observed at 0 eV for Mn-up, Co-dn and Ni-up signifies the metallic character in the respective spin channel. The presence of peaks away from zero energy signifies the absence of energy states at the Fermi level, thereby indicating a semiconducting nature in the Mn-dn, Co-up and Ni-dn spin channels. The variation in the refractive index (n) and extinction coefficients (k) with photon energy replicates the character of the real and imaginary dielectric constants, as shown in Fig. S3 (ESI).† The static refractive index values are 1.47, 1.36 and 1.74 for Mn-up, Co-dn and Ni-up, respectively. The variation in absorption coefficient ( $\alpha$ ) with photon energy is depicted in Fig. 8(A). The critical values of ' $\alpha$ ', beyond which the absorption capabilities of a material increase, illustrate the threshold energies required for electrons to transition from the valance band maxima to the conduction band minima. The transitions mostly from the Cl-p sates of the valance band to the d-states of the M-site in the conduction band give rise to the optical absorption edge. The presence of pseudo gaps in the band structure also gives rise to absorption peaks. It can be seen from the figure that these double perovskites have a wide absorption region. This region is composed of different absorption peaks resulting from various electronic transitions, indicating absorption in the UV-visible region. Moreover, the absorption of photons generates electron-hole pairs, resulting in optical conductivity. The variation in optical conductivity with photon energy is similar to the absorption behaviour. This is due to the fact that absorbed photons generate conduction carriers, and thereby increase conductivity, as

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shown in Fig. 8(B). The peaks of optical conductivity at the static limit of Mn-up, Co-dn and Ni-up reveal their metallic character in the respective channel. In contrast, in the other spin channel, Mn-dn, Co-up and Ni-dn conduct only after the absorption edge, signifying the semiconducting channel. Hence, there results give overall support for the half-metallic character of double perovskites. The total absorption and conductivity of a spin polarized system is the sum of individual channels plus the peaks contributed by inter-channel transitions. We presented the total absorption power and conductivity without considering the inter-channel transitions of the double perovskites in Fig. S4 (ESI).† The peaks in the energy range of 0-6 eV strongly support the capability of these materials to absorb photons in the UV-visible region, confirming their suitability to be employed in various optoelectronic devices.

#### Conclusion

We have successfully established the structural and mechanical stability of halide double perovskites  $\mathrm{Cs_2NaMCl_6}$  (M = Mn, Co and Ni): the results would attract the attention of experimentalists to synthesize these novel materials. Their spin polarized electronic band structure indicates the half-metallic character for these materials. The total magnetic moment and distribution of energy levels signify that  $\mathrm{Mn^{3^+}}$  and  $\mathrm{Co^{3^+}}$  form a high-spin state, while  $\mathrm{Ni^{3^+}}$  is in a low-spin state. The formation of the low-spin state is probably due to the increase in the number of electrons at the central atom. The presence of the integral magnetic moment of  $4\mu_\mathrm{B}$ ,  $4\mu_\mathrm{B}$  and  $1\mu_\mathrm{B}$  for the Mn-, Co- and Ni-based double perovskites, respectively, supports the presence of

half-metallic character. The variation in dielectric constant, absorption coefficient and optical conductivity with energy indicates the absorption of photons in the visible-UV region and also supports the presence of half-metallic character.

## **Authors contribution**

All the calculations have been carried out by Mr Shabir Ahmad Mir. The manuscript was checked and hence modified by Dr Dinesh C. Gupta.

## Data availability statement

The datasets generated during and/or analyzed during the current study would be available from Mr Shabir Ahmad Mir on reasonable request.

## Conflicts of interest

The authors have no conflict of interest.

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