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1. Introduction

Recently, rare-earth (RE)-based scheelite micro/nanostructures have attracted considerable interest from researchers for various optoelectronic applications, including solid-state lasers, fluorescent lamps, optical sensors, scintillators, light-emitting diodes, fiber optic communications, and biological labeling due to their unique features of tuneable emission, narrow bandgap, high quantum yield and high opto-electric conversion, and excellent thermal and chemical stability.^{1–3} Among them are the alkali RE binary molybdates with the generic

A truncated octahedron NaCe(MO_{4})₂ nanostructure: a potential material for blue emission and acetone sensing†

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Scheelite-type NaCe(MO_{4})₂, a promising nanostructure for optoelectronic applications, has been synthesized using a typical hydrothermal technique, and its structural and microstructural properties have been characterized using several microscopic and spectroscopic techniques. We observed intense blue emission from the 5d–4f transitions of Ce³⁺ within CeO₈, while the presence of mono-(CeO₇) and divacancies (CeO₆) of oxygen cause relatively weak emissions at 488 nm, 462 nm and 531 nm. Our study reveals that the use of trisodium citrate during synthesis plays a significant role to tune the formation of $CeO₇$ and $CeO₆$, which in consequence also modify the band edges of the system. The Commission Internationale de l'Eclairage (CIE) coordinates are found to be within the blue region with a correlated color temperature (CCT) of \sim 7854 K, indicating the potential of NaCe(MoO₄)₂ nanostructures for cold solid-state lighting applications. Moreover, the oxygen deficiencies are found to act as active sites for the selective adsorption of acetone, leading to acetone sensing. Hence, NaCe(MoO₄)₂ nanostructures may be potential smart materials for blue-lighting and acetone sensors. Ab initio calculations have been carried out to obtain theoretical insights into the electronic structure of the bare and oxygen-deficient $\text{NaCe}(\text{MoO}_4)$ ₂ and to understand the quiding parameters for acetone sensing. Our calculations also demonstrate that acetone adsorption occurs mostly through the (112) plane. PAPER
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formula $A^{I}Ln^{III}(MoO₄)₂$ (A = alkali metal, Ln = RE lanthanide), in which the Ln^{3+} embedded in the A^+ lattice exhibits a strong 5d-4f transition depending on the symmetry of the unit cell.^{4,5} Extensive research work has been carried out by various researchers, including us, to understand the parameters influencing the emission. As an example, Cheng et al. have investigated the influence of Li/Ag on the luminescence properties of Li_{1-x} Ag_xLu(MoO₄)₂:Eu³⁺,⁶ while Song *et al*. have explored red emission from Eu³⁺-doped AgGd(MoO₄)₂.⁷ The strong integrated emission and higher color purity observed for LiEu(MoO₄)₂ among the AEu(MoO₄)₂ materials (A = Li⁺, Na⁺ and K^+) have been ascribed to its higher, faster charge transfer due to the shortened and more covalent Li–O bond in comparison with those of its Na or K counterpart.⁸ Most of the conventional techniques (e.g., the Czochralski method, solidstate reaction, etc.) lead to agglomeration with high grain size and irregular morphology;⁹ however, recent studies reveal a remarkable correlation between the morphology and emission as the selection rules of optical transition significantly depend on shape, size, exposure facets, dimensionality, hierarchy, etc. To date, abundant efforts have been devoted to generating various scheelite micro/nanostructures using hydrothermal and other techniques to gain an intuitive understanding of

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optical emissions. As a reference, Liu et al. reported the preparation of uniform tetragonal microspindles and nanoplates using the surfactant ethylenediaminetetraacetic acid (EDTA-2Na),¹⁰ while microrods, microcuboids and microflowers have been obtained via a synthesis technique based on the ligand polyvinylpyrrolidone.¹¹ Zheng et al. synthesized a cotton-shaped 3D flower-like morphology using a trisodium citrate assisted method, 12 whilst uniform spherical nanoparticles have been achieved via a cetyltrimethyl ammonium bromide (CTAB) assisted hydrothermal method.¹³ A triclinic \rightarrow tetragonal phase transition was observed by our group during the CTAB-assisted hydrothermal synthesis of NaCe(WO₄₎₂.^{14,15}

In this manuscript, we report for the first time a NaCe- $(Moo_4)_2$ truncated octahedral nanostructure with inherent oxygen mono- and divacancy (V_O and $2V_O$) defects and describe the role of these defects in its photoluminescence. In addition, V_O and $2V_O$ often accumulate at the surface of nanostructures and trap electrons that facilitate gas-sensing properties.¹⁶⁻¹⁹ Although optical studies have been reported in the literature, reports on the gas-sensing of scheelite nano/microstructures are very rare. As an example, Lin et al. reported the V_O -induced gas sensing activity of NaBi(MoO₄)₂ nanomaterials,²⁰ while our recent study demonstrated the NH₃-sensing ability of NaCe $(\mathrm{WO}_4)_2$.¹⁴ We herein also identify acetone-sensing ability in NaCe(MoO₄)₂ nanostructures, with V_O and 2V_O playing crucial roles in determining the sensitivity. To the best of our knowledge, this is the first report on the acetone-sensing ability of NaCe $(Moo_4)_2$. Additionally, as there have been no in-depth theoretical calculations on the electronic structure of $NaCe(MoO₄)₂$, to achieve a better understanding, we have also calculated the *ab initio* band structure of NaCe $(M_0O_4)_2$. The calculations reveal that the Ce $5d_{z}$, $d_{x^2-y^2}$, and d_{yz} orbitals result in blue and green emissions in the presence of V_O and $2V_O$, respectively. We have demonstrated that the electronic excitation involved the $MoO₄$ tetrahedra and that emission takes place within the CeO₈ polyhedra, while MoO₄ \rightarrow CeO₈ charge transfer is mediated through the $\nu_4(\text{F}_2)$ phonon mode. Overall, it may be inferred that the truncated octahedron $NaCe(Moo₄)₂$ nanostructures have strong potential in violet emission and sensing applications.

2. Experimental section

2.1 Materials and synthesis

To prepare $\text{NaCe}(\text{MoO}_4)_2$ nanomaterials through the typical hydrothermal method, 1.00 mmol (0.434 g) of cerium nitrate $[Ce(NO₃)₃·6H₂O,$ Merck, Germany] and trisodium citrate [Na₃Cit-2H₂O, Merck, Germany] were mixed with 60 ml DI water, followed by stirring for 2 h at room temperature. Another aqueous solution of 2.00 mmol (0.484 g) sodium molybdate [Na₂MoO₄·H₂O, Merck, Germany] in 20 ml DI water was added dropwise to the above solution. After 30 minutes of vigorous stirring, the resulting solution turned a pale yellow colour; the solution was then transferred to a Teflon autoclave for hydrothermal reaction at 180 $^{\circ}$ C for 24 h. The final product was

collected by centrifugation, followed by simultaneous washing with DI water and ethanol, overnight drying at 70° C and calcination at 800 \degree C for 5 h in a muffle furnace. The synthesis was repeated using different concentrations of $Na₃Cit·2H₂O$ (0.00, 0.50, 0.78, and 1.00 mmol, i.e., 0.00, 0.147, 0.229, and 0.294 g) to examine its effect on the final product, and the resulting samples were labelled as $NCMO_{0.00}$, $NCMO_{0.50}$, $NCMO_{0.78}$ and $NCMO_{1.00}$, respectively. The reaction procedure is presented schematically in Fig. S1 of the ESI.†

2.2 Characterization and gas-sensing properties

X-Ray diffraction (XRD) patterns were obtained using a Rigaku Ultima III powder diffractometer equipped with CuK_{α} radiation $(\lambda = 1.54056 \text{ Å})$ and utilized to examine the phase purity and other crystallographic information through the Rietveld refinement method using the FullProf program suite. Experimental profiles were fitted with a suitable pseudo-Voigt analytical asymmetric function, while the background was fitted with a fourth-order polynomial function. Field emission scanning electron microscopy (FESEM: Hitachi S – 4800, operated at 5 kV) was used to inspect the morphology of the samples, while a transmission electron microscope (TEM-2100 Plus Electronmicroscope, 200 kV) was used to obtain high-resolution microscopic images and selected area electron diffraction (SAED). Raman spectra obtained using an alpha 300 Witec with a 530 nm laser (power: 3 mW; spot size: $2 \mu m$) and Fourier transform infrared (IR) spectra recorded using an IR Prestige were utilized to examine the short-range structural distortion. X-Ray photoelectron spectra (XPS) were collected using a PHI Versa Probe III Scanning XPS Microprobe with an Al K source. The optical properties of the as-prepared samples were investigated at room temperature using UV-Vis spectroscopy (JASCO V650) and photoluminescence spectroscopy (FP-8300, JASCO, 100 W Xe lamp). Paper More (composition, Article articles and composition of the Diversed on 2022. The main of the Campion of the Campion of the Creative Common Creative Common Creative Common Creative Common Creative Common Creative Comm

Gas-sensing properties were examined using Taguchi-type sensor module, in which a hollow cylindrical Al_2O_3 substrate having Pt electrodes was coated with a thick slurry of the nanostructured NCMO materials through the drop-casting method. The slurry was prepared by mixing the as-prepared samples with isopropyl alcohol as a binder, and the coated substrates were heated at 100 \degree C for 12 h to remove residual solvents. Ni–Cr wire was inserted through the hole of the substrate as a heating element to provide the required temperature by adjusting the applied voltage across the wire. Gases supplied from cylinders, balanced with air, were used during measurement of the sensing performance with an Agilent 34461A digital multimeter interfaced with data logger software.

2.3 Calculation of band structure and investigation of gas-sensing mechanism using ab initio density functional theory

It is well-known that the electronic band structure provides theoretical insight into the opto-electronic properties of any material; hence, we computed the spin-polarized band structure, density of states (DOS), partial DOS (pDOS), total DOS (TDOS) and projected DOS (PDOS) using VASP.²¹ Here, the

plane-wave pseudo-potential (PAW) was used, while the Perdew– Burke–Ernzerhof (PBE) exchange correlation and ultra-soft potential as basis set were considered.²² We employed the valence electrons of the respective atoms as follows: Na atom $(1s^2 2s^2 2p^6)$ 3s¹), Ce atom (5s² 5p⁶ 4f¹ 5d¹ 6s²), Mo atom (4p⁶ 5s² 4d¹⁰), and O atom (2s 2 2p⁴). For this calculation, 5 \times 5 \times 7 Monkhorst-Pack *k* points were used, as this allowed an accurate representation of the Brillouin zone and the electronic properties of the material. Prior to calculating the band structure along $\Gamma \to X \to H_1 \to C \to H \to$ $Y \rightarrow \Gamma$, the structure was optimized with the lowest single-point ground state energy by fixing the cut-off energy at 520 eV, while the convergence was tested between 220 and 620 eV. A maximum atomic displacement of \sim 5 \times 10^{-4} Å and stress of \sim 0.02 GPa were considered for this calculation. Each atom was exposed to a 0.01 eV Hellmann–Feynman force to ensure stable convergence, while the EDIFF and force EDIFG parameters were adjusted to 10^{-6} eV and 10^{-3} eV, respectively, to ensure high-precision computational convergence.¹⁴ Materials Advances

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In order obtain theoretical insight into the influence of the defects on the gas-sensing performance, we calculated the adsorption energies of acetone on the NCMO surface as shown in Table S2 (ESI†). Herein, we have modelled the tetragonal NCMO (112) plane with supercell dimension of 7.573 Å \times 17.823 Å \times 32.422 Å. The k-point mesh for this calculation was optimized at $3 \times 3 \times 1$, while the thickness of the slab and

the vacuum level were \sim 3 nm and 15 Å, respectively. In these calculations, the bottom half of the layers were frozen, whereas the top half of the layers with acetone as the adsorbent were set to relax. The adsorption energy was then calculated using eqn (1):

$$
E_{\rm ads} = E_{\rm NCMO \text{-}acetone} - E_{\rm NCMO} - E_{\rm acetone}
$$
 (1)

where $E_{\text{NCMO}+a cetone}$, E_{NCMO} and E_{acetone} represent the total ground state energies of NCMO in combination with the acetone molecule, bare NCMO and acetone gas, respectively. In order to examine the influence of defects on the gas-sensing activity, we carried out the above calculations by removing one or two oxygen atoms from the (112) plane of NCMO, as shown in Fig. S2 (ESI†).

3. Results and discussion

3.1 Structural and morphological studies using XRD, FESEM and TEM

The XRD patterns of all the samples (Fig. 1) closely match with that of scheelite-type tetragonal NCMO with the space group $I4_1/a$ (C_{4h} ⁶, ICDD Database No. 04-007-5489). The absence of any peak other than those of NCMO confirms the phase purity of the synthesized nanostructures, while the sharp diffraction

Fig. 1 XRD patterns of the NaCe(MoO₄)₂ samples (a) NCMO_{0.00}, (b) NCMO_{0.50}, (c) NCMO_{0.78} and (d) NCMO_{1.00}, respectively.

peaks indicate good crystallinity. The unit cell is represented in Fig. S3, and Table S1 in the ESI† summarizes the structural parameters as obtained from Rietveld refinement, which was carried out until satisfactory convergence of χ^2 , $R_{\rm p}$ and $R_{\rm wp}$ were achieved.²³ As shown in Fig. S3 (ESI†), the unit cell of NCMO comprises Na and Ce atoms at the 4b sites and is coordinated to eight O atoms at 16f site via two different bond lengths to produce $(Na/Ce)O₈$ polyhedra; in turn, these polyhedrons are connected to MoO4 tetrahedra with Mo at the 4a site. The unit cell volume, c value, c/a ratio, Na/Ce-O bond lengths, and the Na/Ce–O–Mo and O–Mo–O bond angles (but not the Mo–O bond lengths) decrease monotonically from $NCMO_{0.00}$ to $NCMO_{1.00}$, indicating an enhancement in the distortion of the unit cells. The $(Na/Ce)O_8$ polyhedral (N) and MoO₄ tetrahedral (K) distortions were calculated using the expressions $N = \frac{\alpha_{1(O-Mo-O)}}{n}$ $\frac{\alpha_{1(O-Mo-O)}\alpha_{2(O-Mo-O)}^{2}}{(\alpha_{2(O-Mo-O)})}$ and $\alpha_{2(O-Mo-O)}$ denote the

two different bond angles, and $K = \frac{d_1(N_{\rm a},C_{\rm c}-O)}{d_1}$ $\frac{d_{1(Na, Ce-O)}}{d_{2(Na, Ce-O)}}$, where $d_{1(Na, Ce-O)}$

and $d_{2(Na, Ce-O)}$ represent the two different bond lengths and were found to be $N = 1.011, 1.029, 1.043, 1.052$ and $K = 1.0587$, 1.0827, 1.0838, 1.0842 for NCMO_{0.00}, NCMO_{0.50}, NCMO_{0.78} and $NCMO_{1.00}$, respectively.^{25,26} According to previous studies, a decrease in the c/a ratio is associated with the partial substitution of Na $^{\rm +}$ by Ce $^{\rm 3+},^{\rm 27,28}$ while the perquisite charge neutrality is balanced by O vacancies (V_O) , causing increases in K and N.

It can be noted from the FESEM image that $NCMO_{0.00}$ possesses an irregular shape (Fig. 2(a)), while a gradual transformation from an irregular to a truncated octahedral shape was observed for $NCMO_{0.50}$, $NCMO_{0.78}$ and $NCMO_{1.00}$ (Fig. 2(b)-(d)) suggesting that $Na₃C$ it has significant role during hydrothermal reaction to tune morphology. Mapping of the samples (Fig. S4, ESI†) clearly reveals even distribution of elements. For a better understanding of the shape of these microstructures, transmission electron microscopy (TEM) was further employed. Fig. 2(e) depicts a typical TEM image of $NCMO_{1.00}$, which corroborates the truncated octahedral shape, while the selected area diffraction pattern (Fig. 2(f)) consists of five diffraction spots corresponding to the (112) , (211) , (123) , (222) and (224) planes of tetragonal NCMO, denoting its crystalline character.

3.2 Investigations of local structure using FTIR, Raman and X-ray photoelectron spectroscopies

To understand the influence of V_O on the short-range structural distortions of $(Na/Ce)O_8$ polyhedra and MoO₄ tetrahedra, we obtained the FTIR and Raman spectra of our synthesized samples. The various peaks in the absorption band (shown in Fig. S5 of the (ESI†)) between 500 and 1000 $\rm cm^{-1}$ are attributed to the various active internal vibrations of $MoO₄$ tetrahedra. Commonly, MoO4 tetrahedra crystallize in either regular $\rm (Mo^RO_4)$ or distorted form $\rm (Mo^DO_4)$ due to $\rm V_O$, while the vibration $(\varGamma_{\rm Td})$ of the isolated Mo $^{\rm R}$ O₄ is represented by:²⁹

$$
\Gamma_{\text{Td}} = \mathbf{A}_1 + \mathbf{E} + 2\mathbf{F}_2 \tag{2}
$$

Fig. 2 FESEM images of NCMO samples (a) $NCMO_{0.00}$, (b) $NCMO_{0.50}$, (c) NCMO_{0.78} and (d) NCMO_{1.00}, (e) and (f) TEM image and SAED pattern of $NCMO_{1.00}$

However, the symmetry of the tetrahedra changes to S_4 in NCMO, for which the IR active modes include $\nu_1(A_1)$, $\nu_2(E_1)$, $\nu_3(F_2)$, $\nu_4(F_2)$, one free rotation $\nu_{f,r}(F_1)$ and one translation F_2 . In contrast to Mo^RO_4 , Mo^DO_4 , which has the C_{2h}^6 space group, exhibits four formula units per crystallographic unit cell with $2Mo(C_2)$, $4Mo(C_1)$, $4Ce(C_1)$ and $24O(C_1)$ symmetries of the atoms, indicating splitting of the vibrational modes (*i.e.*, O \rightarrow $Mo(C_1) \rightarrow O$ and $O \rightarrow Mo(C_2) \rightarrow O$) within Mo^DO_4 due to lower site symmetry. 30 The two bands at 650–749 and 750–1000 $\rm cm^{-1}$ are ascribed to $\nu_3(F_2)$ and $\nu_4(F_2)$, while the peak at 670 cm⁻¹ is assigned to $\nu_2(E_1)$. Careful deconvolution (Fig. S6, ESI†) of the $\nu_3(F_2)$ and $\nu_4(F_2)$ modes yields three peaks (*i.e.*, 693, 705, 722 cm^{-1} and 813, 858, 909 cm^{-1}). As per the bond lengthstretching frequency correlation described by Hardcastle et al., an increase in the bond length decreases the vibrational frequency. 31 Hence, the vibrational mode with lower energy corresponds to $Mo^DO₄$, while the high-energy peaks are ascribed to Mo $^{\rm R}$ O₄. Therefore, the peaks at 722 and 909 $\rm cm^{-1}$ are attributed to the $\nu_3(\mathrm{F}_2)$ and $\nu_4(\mathrm{F}_2)$ modes of Mo^RO₄, respectively, while the peaks at 693/705 $\rm cm^{-1}$ and 813/858 $\rm cm^{-1}$ correspond to the $\nu_3(\rm F_2)$ and ν_4 (F₂) mode of vibration within Mo^DO₄.³⁰ The Mo^DO₄/ $\mathrm{Mo}^{\mathrm{R}}\mathrm{O}_4$ ratio, which was calculated from the areas under the curve, was found to be 0.8, 1.0, 1.6 and 2.9 for $NCMO_{0.00}$, $NCMO_{0.50}$, $NCMO_{0.78}$ and $NCMO_{1.00}$, respectively, indicating an increase in tetrahedral distortion with increasing Na₃Cit concentration, which corroborates the previous structural studies using XRD.

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The Raman spectra, which were obtained up to the range of 1000 cm^{-1} at ambient temperature (Fig. S7, ESI[†]), consist of a peak at 463 cm^{-1} and three bands at 64–231, 255–425 and 681–966 $\rm cm^{-1}$. In general, Raman spectra of scheelite materials at the Γ -point is expressed as:

$$
\Gamma = 3A_g + 5B_g + 5E_g \tag{3}
$$

where the non-degenerate A_{α} and B_{α} modes and the doubly degenerate E_g mode correspond to the various external and internal vibrations of $MoO₄$ and $CeO₈$.²⁹ The peak at 64- 231 cm^{-1} corresponds to the rotation and translation of the MoO₄ tetrahedra and Na⁺ and Ce³⁺, respectively. The sharp peak is attributed to the external vibration of $CeO₈$, which is in good agreement with previous results, 32 while the bands are ascribed to the internal bending of the $MoO₄$ tetrahedra. After thorough examination, the two peaks at 318 and 380 $\rm cm^{-1}$ were assigned to the symmetric A_g bending and antisymmetric B_g bending vibrational modes of $MoO₄$, respectively.³³ While the peak at 463 $\rm cm^{-1}$ is assigned to antisymmetric $\rm A_g$ bending mode of $MoO₄$. The other two Raman peaks at 823 and 889 $cm⁻¹$ are assigned to $(O \rightarrow Mo \rightarrow O)$ antisymmetric stretching B_g and $(O \leftarrow Mo \rightarrow O)$ symmetric A_g mode of vibration.^{29,34,35} The decrease in Raman intensity with increasing Na₃Cit concentration is believed to be related to the increase of V_O and antisite defects, in agreement with the XRD and FTIR studies.^{36,37}

X-Ray photoelectron spectroscopy (XPS), a well-known technique to investigate the chemical state of elements, was adopted here to examine V_O -associated changes in the valence states of Na, Ce, Mo and O, as they are highly sensitive to the local electronic environment. The binding energy data of all elements were adjusted with respect to the binding energy (\sim 284.6 eV) of surface-adsorbed atmospheric C 1s.¹⁴ The survey scans for all the synthesized samples are presented in Fig. S8 of the ESI,† while the high-resolution spectra of Mo, Ce, and O are illustrated in Fig. S9–S11 of the ESI.† Two peaks (i.e., 232.7 and 236.0 eV) of Mo can be readily ascribed to the spin–orbit splitting of the $3d_{5/2}$ and $3d_{3/2}$ orbitals of Mo⁶⁺, while careful deconvolution (Fig. S9(a)–(d), ESI†) reveals the presence of two peaks for the two orbitals (i.e., 232.5 and 232.9 eV for $3d_{5/2}$ and 235.7 and 236.2 eV for $3d_{3/2}$), which were assigned as $Mod_{5/2}(I)$, $Mod_{5/2}(II)$ and $Mod_{3/2}(I)$, $Mod_{3/2}(II)$, respectively. Accordingly, we ascribed Mo3d $_{5/2}$ (11) and Mo3d $_{3/2}$ (11) to Mo $^{\rm R} \rm O_{4},$ while $\text{Mo}3d_{5/2}(i)$ and $\text{Mo}3d_{3/2}(i)$ corresponded to Mo^DO_4 ; their difference can be assigned to the V_O -induced changes in electronic repulsion. In brief, electrons are located on O due to its strong electronegativity, which makes the Mo–O bond significantly ionic, while V_O increases the effective charge on Mo. Hence, the electron–electron repulsion on Mo reduces the binding energies of the Mo $3d_{5/2}$ and $3d_{3/2}$ orbitals in Mo^DO₄. In this context, the smaller energy difference between $3d_{5/2}$ and $3d_{3/2}$ in Mo^DO₄ with respect to Mo^RO₄ (*i.e.*, 3.23 and 3.19 eV) again indicates the influence of the local field generated from $V_{\rm O}$ on spin–orbit splitting. The weighted percentage of $Mo^{\rm D}O_4$ calculated from the area under the curve was found to be \sim 26, 32, 35 and 42% for NCMO_{0.00}, NCMO_{0.50}, NCMO_{0.78} and $NCMO_{1.00}$, respectively, which agrees well with the previous

Fig. 3 Schematic diagram of oxygen vacancies of the $CeO₈$ polyhedral unit.

finding of increasing $Mo^DO₄$ content. As illustrated in Fig. S10(a)–(d) (ESI†), the two asymmetric peaks observed at 877– 891 and 894–912 eV are attributed to the spin–orbit splitting of the $3d_{5/2}$ and $3d_{3/2}$ orbitals of Ce³⁺. Similar to the Mo orbitals, careful deconvolution of the asymmetric $d_{5/2}$ peak yields three peaks at 882.0, 884.6, and 886.9 eV, which were designated as Ce3d_{5/2}(I), Ce3d_{5/2}(II) and Ce3d_{5/2}(III), indicating the presence of Ce with three different oxidation states. $Ce3d_{5/2}(m)$ was assigned to the Ce of the regular CeO_8 polyhedra, while Ce3d_{5/2}(II) and Ce3d_{5/2}(I) correspond to Ce belonging to the $CeO₈$ polyhedra with one or two oxygen vacancies (ca. V_O and $2V_O$) and with the nomenclature CeO₇ and CeO₆, respectively. In this context, it may be stated that $CeO₇$ and $CeO₆$ represent deformed CeO₈ polyhedra with one or two nearby $Mo^DO₄$ and $Mo^DO₄$ subunits (shown schematically in Fig. 3). Due to smaller difference in electronegativity between Ce and O, the Ce–O bond is supposed to be more covalent; therefore, V_O plays the predominant role in terms of this covalence and the binding energy. We observed from the Rietveld analysis that V_O shortens Ce–O bond lengths, which in consequence increases electron–electron repulsion at Ce site; thus, the binding energy of the 3d orbitals is decreased in the distorted polyhedra.³⁸ The analysis showed that the weighted percentage of Ce_a increases (\sim 19, 23, 37, 41%), while there is a significant decrease in the weighted percentage of Ce_b (\sim 48, 46, 37, 26%) from NCMO_{0.00} to NCMO_{1.00}. On the other hand, Ce_c remains almost unchanged for all samples $(\sim$ 30%). Therefore, the analysis indicates greater generation of $2V_O$ at the cost of V_O ; hence, the tendency for V_O cluster formation increases. The XPS of O 1s displays asymmetry, indicating the presence of two different O species. A careful analysis reveals two O 1s peaks (Fig. S11, ESI†) at 530.6 and 532.4 eV, corresponding to lattice O and the O atom of the $Mo^DO₄$ tetrahedra.³⁹ The increasing ratio of the area under the fitted curves illustrates the monotonic increase in $Mo^DO₄$ tetrahedra from $NCMO_{0.00}$ to $NCMO_{1.00}$. Materials Advances

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3.3 Optical properties of the as-prepared NCMO samples using UV-Vis and photoluminescence spectroscopy

The optical band gap $(E_{\rm g})$ was calculated to be \sim 3.07, 3.05, 3.02 and 2.99 eV for NCMO_{0.00}, NCMO_{0.50}, NCMO_{0.78} and NCMO_{1.00}

based on the UV-Vis absorption spectra (shown in Fig. S12, ESI†). This variation can be understood as follows: our band structure calculations (discussed later) reveal that the valence band (VB) is made up of O 2p–Mo 4d hybridization, while the conduction band (CB) comprises Mo 4d, Ce 4f and a minute contribution from O 2p orbitals. The decrease of Mo–O bond length in the presence of V_O and $2V_O$ reduces the O 2p–Mo 4d overlap, which causes a blue-shift of the VB, and subsequently, E_g is reduced. All the samples exhibit a broad peak at 380 nm in their photoluminescence excitation (PLE) spectra, which were obtained using an emission wavelength λ_{em} of 531 nm (Fig. S13, ESI†), while the emission spectra recorded in the visible region using an excitation wavelength of $\lambda_{\rm ex}$ = 380 nm are depicted in Fig. $4(a)$ – (d) . The PLE peak was assigned to

O 2p–Mo 4d charge transfer absorption within $MO₄$ tetrahedra.40,41 The emission spectra have four distinct peaks in the blue (i.e. blue I, blue II, blue III) and green regions, with an intense blue I peak being observed at 440 nm (22727 cm^{-1}) , while relatively weaker blue II, blue III peaks and a prominent green peak are observed at 462 nm (21 645 $\rm cm^{-1})$, 488 nm $(20 492 cm^{-1})$ and 531 nm $(18 832 cm^{-1})$, respectively. The Stokes shift of the excitation and emission spectra were noted to be \sim 3588 cm $^{-1}$, suggesting the absence of reabsorption and interference among the various emissions. 42 Although understanding the emission spectra of lanthanides is difficult for several reasons, including symmetry, the crystal field surrounding the host crystal, anion polarizability, and covalence of the host crystal, the present emissions are believed to be 5d–4f

Fig. 4 Photoluminescence spectra of (a) NCMO_{0.00}, (b) NCMO_{0.50}, (c) NCMO_{0.78} and (d) NCMO_{1.00}. (e) Normalized PL intensity ratio of emission wavelengths. (f) CIE chromaticity diagram of the NCMO phosphors.

transitions within Ce^{3+43} and can be validated as follows. The emission wavelength (λ in nm) for intra-5d-4f transitions can be calculated using eqn (4) : $44,45$

$$
\frac{1}{\lambda} = \frac{Q^*}{hc} \left[1 - \left(\frac{V}{4}\right)^{\frac{1}{V}} 10^{-\frac{(nrE_a)}{80}} \right]
$$
(4)

where, Q^* , V , 'n', E_a and 'r' represent the energy of the 5d band edge of a free Ce³⁺ ion (= 50 000 $\rm cm^{-1}$), the valence of the Ce³⁺, the number of anions in the immediate shell around the Ce^{3+} , the electron affinity of the atoms forming the anions $(\sim 2.16 \text{ eV})$ and the difference between the average bond length and radius of Ce^{3+} within the CeO_8 polyhedra (1.03 Å). Very careful calculation yields $\lambda \sim 449$ nm, which corresponds to blue I emission. Previous studies indicate that V_O and $2V_O$ can significantly modify the 5d–4f transitions of lanthanide atoms, resulting in different emissions. As an example, Sokolenko *et al.*⁴⁶ assigned green-red emission to $WO_3 \cdot V_O^{\bullet\bullet}$ oxygendeficient complexes, while Korzhik et al. ascribed green emission to WO₃ centres.⁴⁷ Considering CeO₇ and CeO₆ to be V_Oand $2V_{\Omega}$ -containing distorted CeO₈ polyhedra, we have calculated λ values of 487 and 537 nm for intra-Ce³⁺ 5d-4f transitions using eqn (4), which are in good agreement with the blue III and green emissions (schematically represented in Fig. S14, ESI†). Materials Advances

transitions within x^{p+2} and each by suidated as follows. The the CP using eqn (7)^{23,53}

the calculated unit per literations As 4 If transitions each
 $\frac{1}{2} = \frac{Q_p}{R} \left[1 - \left(\frac{p}{R}\right)^{\frac{1}{2}} \frac{1$

The shift of the centroids of the d-orbitals (in eV) of Ce^{3+} , $\varepsilon_c(1,3^+,Ce^{3^+})$ in CeO₇ and CeO₆ are believed to tune the 5d-4f transition, giving various emissions (see discussion in the ESI†). Careful monitoring of the green emission reveals the presence of a phonon side band (PSB) at 516 nm with a phonon energy of 930 cm^{-1} , which exactly matches the energy of the $\nu_4(F_2)$ phonon mode, indicating that MoO₄ \rightarrow CeO₆ charge transfer is mediated by the $\nu_4(F_2)$ phonon mode. In this context, the Huang–Rhys (S) factor, an indicator of electron–phonon coupling, was calculated using eqn (5) and (6) : $48,49$

$$
S = \frac{I_{PSB}}{I_{ZP}}\tag{5}
$$

$$
\frac{I_{\rm IP}}{I_{\rm ZP}} = \frac{W_1(S, \langle m \rangle)}{W_0(S, \langle m \rangle)} = \begin{cases} S\langle 1 + m \rangle, p \ge 0\\ S\langle m \rangle, p < 0 \end{cases} \tag{6}
$$

where, I_{PSB} , I_{ZP} and I_{1P} are the integrated intensities of the PSB, zero-phonon line and one-phonon line. As this emission includes a single phonon ($p \geq 0$ in eqn (6)), and considering the ${}^{5}D_{0}$ ⁻²F_{5/2} transition to be purely dipolar, we may write $I_{\rm PSB}$ = I_{1P} . In addition, $\langle 1 + m \rangle$ can also be considered to be 1 due to the higher phonon energy at room temperature. The decrease in the S-factor (0.098, 0.077, 0.032 and 0.039 for the four respective samples) is ascribed to increasing distortion in the lattice structure due to V_{Ω} and $2V_{\Omega}$.

The Commission International De l'Eclairage (CIE) coordinates (Fig. 4(f)), which were calculated to be (0.180, 0.149), $(0.180, 0.147), (0.170, 158)$ and $(0.172, 0.148)$ for the NCMO_{0.00}, $NCMO_{0.50}$, $NCMO_{0.78}$ and $NCMO_{1.00}$ samples, are well spread throughout the blue region of the visible spectrum, indicating the color purity (CP) of the NCMO samples.⁵⁰ We also validated the CP using eqn (7) :^{51,52}

$$
CP = \frac{\sqrt{(x - x_i)^2 + (y - y_i)^2}}{\sqrt{(x_d - x_i)^2 + (y_d - y_i)^2}} \times 100\%
$$
 (7)

where (x, y) denotes the color coordinates of the phosphor; (x_i, y_i) is the 1931 CIE Standard Source's illuminant point, with color coordinates of (0.3101, 0.3162); and (x_d, y_d) is the color coordinates of the dominant wavelength. Outstanding CP values of \sim 90–94% were observed for the samples, indicating that NCMO could be a good candidate for blue-light-emitting applications. The correlated colour temperature (CCT) was calculated using McCamy's relation:⁵³

$$
CCT = 449n3 + 3525n2 + 6823.3n + 5520.3
$$
 (8)

where $n = \frac{(x - 0.3320)}{(0.1858 \cdot x)}$ $\frac{(0.1858 - y)}{(0.1858 - y)}$ and (x, y) represent the chromaticity coordinates. The CCT values were \sim 7819 K, 7630 K, 7715 K and 7854 K for the samples $NCMO_{0.00}$, $NCMO_{0.50}$, $NCMO_{0.78}$ and NCMO_{1.00}, respectively. The high CCT indicates that NCMO can be used for cold blue lighting.

3.4 Investigations of gas-sensing properties of NCMO nanostructures

The gas-sensing abilities of metal oxide nanostructures involve a solid–gas interfacial reaction mechanism in which the adsorption/desorption of the targeted gas molecules on the surface of the nanostructure leads to significant changes in resistance. A careful literature survey indicates that the following redox reactions between the nanostructure and acetone (CH_3COCH_3) , ethanol $(C₂H₅OH)$, ammonia (NH₃), methanol (CH₃OH), formaldehyde (HCHO) take place in the presence of O_2 ⁻(ads).

$$
CH_{3}COCH_{3}(gas) + 8O_{2}^{-}(ads) \leftrightarrow 3H_{2}O(gas) + 3CO_{2}(gas) + 8e^{-}
$$
\n(9)
\n
$$
CH_{3}CH_{2}OH(gas) + O_{2}^{-}(ads) \leftrightarrow 2H_{2}O(gas) + C_{2}H_{2}O + 2e^{-}
$$
\n(10)
\n
$$
4NH_{3}(gas) + 3O_{2}^{-}(ads) \leftrightarrow 6H_{2}O(gas) + 2N_{2}(gas) + 3e^{-}
$$
\n(11)
\n
$$
CH_{3}OH(gas) + O_{2}^{-}(ads) \leftrightarrow 2H_{2}O(gas) + CO(gas) + 2e^{-}
$$
\n(12)
\n
$$
HCHO(mse) + O_{2}^{-}(ads) \leftrightarrow H_{2}O(mse) + CO_{2}(mse) + 2e^{-}
$$

$$
HCHO(gas) + O_2^-(ads) \leftrightarrow H_2O(gas) + CO_2(gas) + 2e^-
$$
\n(13)

Our previous investigations indicate that V_O and $2V_O$ defects are often ionized (e.g., $CeO_7 \cdot V_O^{\bullet}$, $CeO_6 \cdot 2V_O^{\bullet}$) and act as electron donors ([CeO $_8$]' and [CeO $_8$]"), which can be understood from Kröger-Vink notation:

$$
[CeO_8]^x + [CeO_7 \cdot V_O^x] \rightarrow [CeO_8]^{\prime} + [CeO_7 \cdot V_O^{\bullet}] \qquad \quad \textbf{(14)}
$$

$$
[CeO_8]^x + [CeO_6 \cdot 2V_O^x] \rightarrow [CeO_8]^{"} + [CeO_6 \cdot 2V_O^{\bullet}]\qquad (15)
$$

These $[CeO₈]'$ and $[CeO₈]'$ facilitate electron transfer from the host materials to the chemisorbed oxygen molecule to generate $O_2^-(ads)$, as described by the reaction: $O_2(gas) + e^- \leftrightarrow O_2^-(ads).$ ^{54–57}

Fig. 5 (a) Temperature calibration curves of NCMO_{0.00}, NCMO_{0.50}, NCMO_{0.78} and NCMO_{1.00}, respectively. (b) Selectivity ofNCMO_{0.00}, NCMO_{0.50}, NCMO_{0.78} and NCMO_{1.00} sensors measured at an operating temperature of 300 °C. (c) Linear fitting of the logarithm of sensitivity and logarithm of acetone concentration data of all samples. (d) Dynamic acetone response curves of the NCMO_{1.00} sensor to acetone concentrations from 1-100 ppm at 300 °C. (e) Response and recovery time of the NCMO_{1.00}-based sensor for 1 ppm acetone at 300 °C. (f) Long-term stability of the NCMO_{1.00}-based sensor in sensing 1 ppm acetone measured over 90 days.

In order to validate the gas-sensing ability of our synthesized NCMO nanostructures as a proof of concept, we carried out a series of sensing measurements using $NCMO_{0.00}$, $NCMO_{0.50}$, $NCMO_{0.78}$ and $NCMO_{1.00}$ as a resistive sensor probe against $NH₃$, $CH₃COCH₃$, $C₂H₅OH$, CH₃OH and HCHO at different operating temperatures (200–450 \degree C) at intervals of 50 \degree C. An optimum operating temperature of \sim 300 °C was observed for all the samples (shown in Fig. 5(a)), and they are highly selective toward $CH₃COCH₃$ (Fig. 5(b), 5.0 ppm concentration) in comparison with

C₂H₅OH, NH₃, CH₃OH, and HCHO $\left(\frac{R_{\rm gas}}{R_{\rm air}} < 1.2\right)$ \sqrt{p} \sqrt{p} . The sensitivity monotonically increases from $NCMO_{0.00}$ to $NCMO_{1.00}$, with the highest sensitivity being observed for $NCMO_{1.00}$. To understand the role of V_O , $2V_O$ in CH_3COCH_3 adsorption/desorption, we fitted the response data using the Freundlich adsorption isotherm as given in eqn (16) and (17).

$$
S_g = 1 + aC_g^b \tag{16}
$$

i.e.,

$$
\log(S_{\rm g}-1) = a + b \log(C_{\rm g}) \tag{17}
$$

where $S_{\rm g}$ and $C_{\rm g}$ represent sensitivity and concentration respectively, while 'a' and 'b' are constants that depend on the electrical charge of the species and stoichiometry of the reactions on the surface of the sensing probe. In the literature, 'b' is noted to be dependent on the adsorbed O species, with $b \leq 0.5$ for O^{2-} adsorption and \approx 1.0 for O^- adsorption.⁵⁸ From slope of the linear regressions (Fig. 5(c)) curves obtained using eqn (17), we calculated *b* values of ~ 0.423 , 0.279, 0.088, 0.053 for NCMO_{1.00}, NCMO_{0.78}, NCMO_{0.50} and NCMO_{0.00}, respectively, suggesting that $2V_O$ acts as a more-active site for O^{2-} generation, facilitating CH_3COCH_3 adsorption. The dynamic sensing responses of $NCMO_{1.00}$ (Fig. 5(d)) measured at 300 °C demonstrate sensitivity values of \sim 1.86, 2.81, 4.95, 6.21 and 6.89 in the presence of 1.0, 5.0, 10.0, 50.0 and 100.0 ppm of $CH₃COCH₃$, while the response and recovery (at 5.0 ppm of CH_3COCH_3 , see Fig. S15(b) of the ESI†) remain the same for six consecutive cycles (error \leq 2%), indicating the excellent reproducibility and reliability of $NCMO_{1.00}$ as a sensor probe in comparison with the other samples (Fig. S15(c–e), ESI†). In addition, the very low response and recovery time of \sim 1.5 s/33.8 s (Fig. 5(e), measured at 10.0 ppm) indicate the fast response of $NCMO_{1.00}$, while its long-term stability was checked and verified over 90 days at intervals of 10 days (Fig. 5(f)).

3.5 Density functional theory (DFT) calculations of NCMO

In order to validate the V_O - and $2V_O$ -associated optical emissions and sensing activity, we calculated the electronic band structure, total density of states (TDOS), and angular momentum projected partial density of states (PDOS) using ab initio density functional theory (DFT), which provides valuable insights to explain the luminescence and gas sensing behaviour. For this, we calculated band structure along several high symmetry k-points within the Brillouin zone. In this calculation, we removed one or two O atoms from the NCMO unit cell to mimic CeO₇ (V_O) and CeO₆ (2V_O). Prior to calculating the electronic band structure, we optimized the unit cell of pure NCMO and obtained lattice parameters $a = b = 5.355$ and

 $c = 11.611$ Å and Na/Ce–O and Mo–O lengths of \sim 2.453 and 1.806 Å, respectively. These were in good agreement with the experimental results, proving the accuracy of our calculations, specifically, the choice of exchange correlation and pseudopotential. The top of the valence band maxima (VBM) was set to zero as a reference for other calculations. As shown in Fig. 6(a), NCMO appears to be an indirect bandgap material ($E_g \sim$ 3.23 eV) with a spin-unpolarized VBM and spin-polarized conduction band minima (CBM) at Γ - and X-points, respectively. The curvature of the VBM is observed to be low, indicating a comparatively high effective hole mass, while the CBM has a high curvature suggesting a low effective electron mass. The TDOS (shown in Fig. 6(b)) and PDOS contributions of O, Mo, Ce and Na (shown in Fig. $S16(c)$ –(f), ESI[†]) demonstrate that the upper part of the VB is primarily composed of the O $2p_x$, $2p_y$, and $2p_z$ orbitals with very small contributions from Mo $2p_x$, $2p_y$, and $2p_z$, giving a low curvature. The lower part of the VB originates from predominant hybridization between Mo $4d_{xy}$, $4d_{xz}$, $4d_{z^2}$ and O $2p_x$, $2p_y$, and $2p_z$ orbitals, whereas $4d_{x^2}$ and $4d_{yz}$ make insignificant contributions. We noted almost no contribution to the VB from the Ce 4f and Na 3s orbitals. The CBM comprises spin-polarized (*i.e.*, up-spin) Ce $4f_{z3}$, $4f_{xz2}$ and $4f_{y3x2}$ orbitals, among which the most predominant contribution comes from $4f_{xz^2}$, hybridized with O $2p_x$, $2p_y$, and $2p_z$ orbitals, giving a higher curvature in comparison with the VBM, while the upper portion of the CB is made up of Mo–4 d_{γ^2} and $4d_{\gamma^2}$. Thus, it may be stated that the $MoO₄$ tetrahedra mostly construct the VB, while the CB is predominantly contributed by the CeO_8 polyhedra, and the band-to-band transition includes a Mo \leftrightarrow Ce charge transfer process. Materials Advances
 $\tkappa = 1.24 \text{ B log}(G_2)$
 $\log 8x_2 - 11 - a + b \log(G_2)$

Where $\frac{8}{3}$ and G_2 expresent essention-properties and on the common and expectation-there are the positive Common and the common and the common an

The band structures, TDOS and PDOS of the V_O^{\bullet} - and $2V_O^{\bullet}$ -containing NCMO samples, designated as NCMO - V_O^{\bullet} and NCMO $-2V_0^{\bullet}$, are represented in Fig. S17 and S18 (ESI†), respectively. Notably, both the VBM and CBM (shown in Fig. S17(a), ESI†) of NCMO – V_O^{\bullet} remain unchanged, while in case of NCMO – $2V_O^{\bullet}$, the CBM band has shifted into the *Γ*-point (shown in Fig. S17(a), ESI†), indicating the transformation from an indirect to a direct E_g nature for NCMO. We observed an $E_{\rm g}$ of 3.01 and 2.95 eV for NCMO – $V_{\rm O}^{\bullet}$ and NCMO – $2V_{\rm O}^{\bullet}$, suggesting a decrease in $E_{\rm g}$ due to $V_{\rm O}^{\bullet}$ and $2V_{\rm O}^{\bullet}$. Thus, the

Fig. 6 (a) Band structure and (b) spin-polarised TDOS of NCMO.

monotonic decrease in the experimentally measured E_g from $NCMO_{0.00}$ to $NCMO_{1.00}$ is believed to be due to increasing V_O and 2V_O, corroborating previous studies. The reduction of E_g can be understood as follows: TDOS (Fig. S17(b) and S18(b), ESI[†]) and PDOS (Fig. S17(c)–(f) and S18(c)–(f), ESI[†]) calculations show that the VB width decreases in NCMO – $2V_O^{\bullet}$ due to the greater localization of the O 2p orbitals, while the curvature increases in NCMO – V_O^{\bullet} and NCMO – $2V_O^{\bullet}$, which is attributed to the enhanced overlap between the Ce $4f_{zx^2}$, $4f_{xz^2}$, $4f_{yz}$, and $4f_{xyz}$ and O $2p_x$, $2p_y$, and $2p_z$ orbitals due to the decrease in the Ce–O bond length in the CeO_8 polyhedra, indicating higher carrier mobility. Further analysis of the TDOS and PDOS reveals that the contributions from Ce $5d_{xy}$ and $5d_{xz}$ to the CB increase in NCMO – V_{O}^{\bullet} and NCMO – $2V_{O}^{\bullet}$. The defect state at 2.44 eV above the valence band in NCMO – V_O^{\bullet} was assigned to Ce $5d_{z^2}$, whilst these defects are found at 2.26 and 2.40 eV above in NCMO – $2V_{O}^{\bullet}$ due to Ce 5d_{z^{2}} and Ce 5d_x², respectively. There-</sub> fore, it can be stated that the blue III emission originates from Ce $5d_{z^2} \rightarrow 4f$ transitions in the CeO₇ polyhedra, while the blue II and green emissions are attributed to Ce $5d_{z} \rightarrow 4f$ and Ce $5d_{x^2} \rightarrow 4f$ transitions in the CeO₆ polyhedra, respectively, as schematically presented in Fig. 7. Paper Montonic decrease in the copies
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From the calculation of the adsorption energies on the (112) plane with the lowest energy for the NCMO, NCMO – V_O^{\bullet} and $NCMO - 2V_O^{\bullet}$ systems using eqn (1), we found the energies listed in Table S2 (ESI†). The E_{ads} (eV) values represent the adsorption energies of the pristine NCMO, NCMO - V_O^{\bullet} , NCMO – $2V_{\text{o}}^{\bullet}$ systems, and it was found that $E_{\text{ads}} < 0$, indicating that the adsorption processes involved in these systems are thermodynamically stable and exothermic, meaning that they release energy during the adsorption process. Additionally, the increasingly negative values of NCMO – V_O^{\bullet} and NCMO – $2V_O^{\bullet}$ suggest the presence of active sites (oxygen vacancies) on the NCMO surface, which create localized electron-rich sites enabling strong interactions to occur between the $CH₃COCH₃$ and NCMO surfaces.⁵⁹ Hence, the highest sensitivity of $NCMO_{1.00}$ (Fig. 5) can purely be assigned to the presence of

Fig. 7 Schematic picture of luminescent transitions within different CeO_8 polyhedra involving different orbitals.

the highest $2V_O$ content, which is in good agreement with the experimental results and theoretical calculations.

4. Conclusion

In summary, this study reports the synthesis and characterization of oxygen-deficient NCMO-truncated octahedral nanostructures, which exhibit intense blue I and green emissions and comparatively low-intensity blue II and blue III emissions. Excitation involves $MoO₄$ tetrahedra, while emissions are attributed to the 5d \rightarrow 4f transitions of Ce in different CeO₈ polyhedral configurations. This study also identified the acetone-sensing ability of NCMO, which has been attributed to $CeO₆$. The valence and conduction bands of NCMO comprise O 2p and O 2p and Ce 5d orbitals, respectively; hence, they are highly sensitive to oxygen vacancies and have a potential role in luminescence and gas sensing. Our study suggests that NCMO may be a promising material for blue-light-emitting and acetone-sensing applications.

Conflicts of interest

There are no conflicts to declare.

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