Electronic structure and luminescence assets in white-light emitting Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ pyro-vanadates: X-ray absorption spectroscopy investigations

Aditya Sharma, Mayora Varshney, Keun-Hwa Chae and Sung Ok Won*

Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$, and Ba$_2$V$_2$O$_7$ pyro-vanadates were synthesized using a modified chemical precipitation method and annealing. Detailed crystal structure, morphology, electronic structure and optical properties were investigated by XRD, UV-visible absorption, FTIR, Raman, FE-SEM, XANES, and photoluminescence spectroscopy. Rietveld refinement on the XRD patterns of Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$, and Ba$_2$V$_2$O$_7$ has confirmed the triclinic structure (space group; $P\overline{1}(2)$) of the pyro-vanadates. The band gap energy of Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$, and Ba$_2$V$_2$O$_7$ is estimated to be $\sim$2.67 eV, $\sim$2.97 eV and $\sim$3.09 eV, respectively. XANES spectra at the Ca L-edge, Sr K-edge and Ba L-edge have confirmed the Ca$^{2+}$, Sr$^{2+}$ and Ba$^{2+}$ ions in the Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ compounds, respectively. V K-edge XANES spectra have strengthened the presence of sub-pentavalent V ions in all of the pyro-vanadates. O K-edge XANES spectra of Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ have shown dominating tetrahedral symmetry of the V ions which is also corroborated with the V K-edge XANES. Broad-band emission spectra, ranging from 400 nm to 700 nm, have been observed from the charge-transfer transitions of VO$_4$ tetrahedra. $^3T_1 \rightarrow ^1A_1$ and $^3T_2 \rightarrow ^1A_1$ transitions, from the VO$_4$ tetrahedra, have provided two distinct emission peaks from the compounds which exhibit a red-shift with the decreasing ionic-radii of alkali-earth metal ions. The mixed compounds, with equal weight proportions, have shown remarkable emission characteristics towards the realization of rare-earth element free white-light-emitting devices.

Introduction

Analogous to the transition metal based pyro-vanadates, M$_2$V$_2$O$_7$ (M = transition metal; Mn, Co, Ni and Cu etc.), the alkali earth metal based A$_2$V$_2$O$_7$ (A = Ca, Sr and Ba etc.) oxides are of special interest because of their promising applications in rare earth element free white-light-emitting devices$^{6-10}$ and microwave-dielectrics.$^{11,12}$

The electron correlation effects (because of the V–O4, V–O5 and V–O6 kinds of conceivably polyhedral configuration of V and O ions) make A$_2$V$_2$O$_7$ oxides attractive candidates for probing their intriguing correlations among the crystal structure, electronic/atomic structure and diverse applications. The alkali earth metal based pyro-vanadates (i.e., Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ etc.) obey the triclinic crystal structure. Their band structure, near the Fermi level, is characteristically formed by the hybridization of the frontier V d-orbitals and O 2p-orbitals. The band structure of the conduction band is shaped by the anti-bonding O 2p and V 3d orbitals and the top of the valence band mainly consists of non-bonding O 2p-orbitals.$^{9,10}$ The orbitals of the alkali earth elements provide a large conduction band-width and do not contribute to the band structure near the Fermi level. The energy-band structure and, thus, the electronic/ optical/ magnetic properties of A$_2$V$_2$O$_7$ oxides are not only sensitive to the amalgamation of foreign elements$^{11-15}$ but are also expected to be modified by the distortion in the V–O polyhedra. A marginal distortion in the metal–oxygen polyhedra can affect the hybridization between the O 2p and metal d orbitals, which indeed regulate the band structure near the Fermi level and, consequently, the electronic/optical properties of the compounds.$^{10,17}$

Optical properties and illumination performance of the compounds are categorically determined by the electronic structure around the stimulated ions and energy transfer processes among them.$^{18-22}$ In case of vanadium-based optical materials, the (VO$_4$)$^{1-}$ groups obey broad and strong charge-transfer (CT) absorption bands near the UV region and, thus, produce broad-emission spectra related to the electronic/atomic structure of the compound where the V ions are coordinated by the four O ligands.$^{6,10,19,20}$ In previous reports, the Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ have shown broad emission characteristics ranging from 400 nm to 700 nm and have signified the concept of vanadium-based and rare-earth-free white-light emitting phosphor materials with high efficiency.$^{6,10,19,20}$ However, synthesis of single phase A$_2$V$_2$O$_7$ oxides is challenging.
because of unpredictable nature of V ions to form secondary phases of VO₂, V₂O₅ etc. under the chemical and/or thermal treatment and, therefore, high-purity phase fabrication of A₂V₂O₇ compounds is still a thirsty area of research for establishing the financially viable and straightforward synthesis approaches.

Besides the single-phase synthesis of A₂V₂O₇ compounds, investigations on the crystal structure and electronic structure properties, which may have diverse V-O hybridization and complex electronic correlation, are also indispensable using the modern, non-destructive and element specific techniques. In this regard, X-ray photoelectron spectroscopy (XPS) had been applied to probe the electronic structure and chemical environment in pyro-vanadates. The XPS is sensitive to a few surface layers of the samples and an insulating/less-conducting sample undergoes severe charging effects during the measurements. Therefore, XPS is not ideally suited to correlate the structural features to the electronic/atomic structure properties of bulk of the material. In this study, in order to investigate the relationship among the crystal structure, orbital hybridization and optical properties of A₂V₂O₇ pyro-vanadates, we have systematically applied the X-ray absorption near edge structure (XANES) spectroscopy technique at O K-edge, V L-edge, Ca L-edge, Sr K-edge, V K-edge and Ba L-edge. Besides providing the detailed experimental results on the absorption edges of constituent elements of the samples, the XANES investigations presented below address the several aspects. First, to specifically look into the dissimilarities in the hybridization of frontier orbitals of V and O ions in Ca₂V₂O₇, Sr₂V₂O₇ and Ba₂V₂O₇, the low energy XANES spectra at O K-edge and V L-edge were studied. Secondly, to understand valence state of constituent elements in the samples, systematic, XANES spectra at Ca L-edge, Ba L-edge, Sr K-edge, V K-edge were collected and analyzed.

**Results**

(a) XRD, UV-visible absorption, FTIR, Raman and FE-SEM studies

Fig. 1 shows the Rietveld refined XRD patterns of pyro-vanadate ceramics sintered at 800 °C for 4 hours. All ceramics flawlessly matched with their relevant JCPDF files: #97-002-0609, #97-007-2200 and #97-003-4320 for Ca₂V₂O₇, Sr₂V₂O₇ and Ba₂V₂O₇, respectively. The initial structure models in the Rietveld refinements of a pyro-vanadate compound were taken from the respective JCPDF files. The insets of Fig. 1 show the magnified view of the simulated patterns. The crystallographic data of Ca₂V₂O₇, Sr₂V₂O₇, and Ba₂V₂O₇ is presented in Table 1. It is noticeable from the Table 1 that all three ceramics have a triclinic structure with P1(2) space group symmetry without forming any secondary phase and, thus, are signifying the single phase formation of pyro-vanadate in this study. Moreover, the lattice parameters and cell volume are higher for the A₂V₂O₇ ceramics with increasing the atomic number of the alkali earth element and are consistent with the previous reports.

Fig. 2(a) shows the UV-visible absorption spectra of Ca₂V₂O₇, Sr₂V₂O₇, and Ba₂V₂O₇ samples. It is noticeable from the Fig. 2 that the Ca₂V₂O₇, Sr₂V₂O₇, and Ba₂V₂O₇ show a sharp absorption edge ~465 nm, ~418 nm and ~402 nm, respectively. Fig. 2(b) shows the band-gap energy estimation from the UV-visible absorption spectra. The band-gap energy was estimated by plotting the hν versus (αhν)² curves (Tauc’s procedure, where α is the absorption coefficient and hν is the photon energy).
energy) and marking intersect at the energy scale. This gives band gap energy of 2.67 eV, 2.97 eV and 3.09 eV for the Ca2V2O7, Sr2V2O7, and Ba2V2O7, respectively. Thus the estimated band-gap energy of the compounds is comparable to the earlier reports.\textsuperscript{18,23,24}

Fig. 2(c) shows the FTIR spectra of the Ca2V2O7, Sr2V2O7, and Ba2V2O7 samples. All of the three samples show few dominating bands \( v1 \) (\(~950–900 \text{ cm}^{-1}\)), \( v2 \) (830–810 \text{ cm}^{-1}), \( v3 \) (750–770 \text{ cm}^{-1}) and \( v4 \) (650–700 \text{ cm}^{-1}). These bands are mainly due to the stretching vibrations of metal-oxygen bonds and bending vibrations of VO\(_4\)\(^{1-}\) groups in triclinic structured A\(_2\)V\(_2\)O\(_7\) pyrovanadates.\textsuperscript{25,26} The \( v4 \) and \( v3 \) bands are the results of V–O–V bridge vibration and asymmetric stretching of longer V–O bond.\textsuperscript{31,32} The rest of the bands (\( v1 \) and \( v2 \)) can be ascribed to symmetric stretching of the other shorter V–O bonds in the samples.\textsuperscript{31,32} Therefore, the feature-rich FTIR spectra convey the existence of VO\(_4\)\(^{1-}\) groups in all of the pyrovanadates.

Fig. 2(d) shows the Raman spectra of Ca2V2O7, Sr2V2O7, and Ba2V2O7 samples. Similar to the FTIR spectra, the Raman spectra have shown several vibration bands of V–O bonds. In all three samples, the Raman modes can be seen within the three distinct regions. The first region is between the 950–800 \text{ cm}^{-1} and can be ascribed to the symmetric stretching vibrations of end VO\(_3\) groups.\textsuperscript{33,34} The second region is from 800 \text{ cm}^{-1} to 400 \text{ cm}^{-1} and attributed to stretching vibrations of bridging V–O–V band bending vibrations.\textsuperscript{35,36} The third region contains bands below the 400 \text{ cm}^{-1} and may be due to the peripheral modes.\textsuperscript{33,34} Though the pyro-vanadates give their rich Raman bands due to the V–O or V–O–V bonds in the samples, however, their position is sensitive to the type of alkali metal element and defect chemistry.\textsuperscript{31,33}

To investigate the surface morphology of the Ca2V2O7, Sr2V2O7, and Ba2V2O7 samples, systematic, FE-SEM images were collected and are presented in Fig. 3(a–c). It is noticeable from the micrographs that all pyro-vanadate have unusual and agglomerated morphology. This unusual morphology may arise due to the diffusion of particles during annealing of samples.\textsuperscript{26,27} The FE-SEM images ruled out the formation of fascinating morphologies (i.e., nano-rod, nano-wire or other structures) and, therefore, the observed luminescence properties are expected to be related only to the electronic/atomic structure properties but not due to the diverse surface/interface related phenomena in nanostructures.

![Experimental and Rietveld refined XRD patterns of Ca2V2O7, Sr2V2O7, and Ba2V2O7. The insets show the magnified view of simulated and experimental data.](image)

**Fig. 1** Experimental and Rietveld refined XRD patterns of Ca2V2O7, Sr2V2O7, and Ba2V2O7. The insets show the magnified view of simulated and experimental data.

**Table 1** Refinement parameters and crystallographic data of Ca2V2O7, Sr2V2O7, and Ba2V2O7. Numbers in parenthesis show error therein

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![UV-visible absorption spectra of Ca2V2O7, Sr2V2O7, and Ba2V2O7.](image)

**Fig. 2** (a) UV-visible absorption spectra of Ca2V2O7, Sr2V2O7, and Ba2V2O7. (b) Shows the Tauck’s plot for determining the band-gap energy. (c) FTIR spectra. (d) Raman spectra of the samples.
Electronic structure study by XANES

Formation of secondary phases in the host, lattice defects and variation in the coordination chemistry of the elements may facilitate ambiguous changes in the valence state of the metal ions in the pyro-vanadates. Therefore, to understand the valence state of Ca, Sr and Ba elements in the samples XANES spectra at Ca L3,2-edge, Sr K-edge and Ba L 3,2-edge were collected and the normalized spectra, along with first derivative spectra, are presented in the Fig. 4 (a–c). The reference samples of CaO, SrO, and BaO, in which valence state of alkali earth cations is 2+ (i.e., Ca2+, Sr2+ and Ba2+), were also scanned under the similar experimental conditions and the spectra are plotted along with the A2V2O7 samples. It is noticeable from Fig. 4 that the edge-energy position and spectral features of reference CaO, SrO and BaO samples fairly resemble with the Ca2V2O7, Sr2V2O7, and Ba2V2O7, respectively, and are strengthening the 2+ valence state of the alkali earth cations in A2V2O7 pyro-vanadates of this study. Interestingly, Ba L2,3-edge XANES of Ba2V2O7 sample shows additional spectral features between the 5400 eV to 5600 eV. V K-edge spectral features originate within this energy range. In the present study, the existence of V ions in Ba2V2O7 sample is responsible for such spectral features. We have not seen any ambiguous spectral feature of secondary phases and, thus, the XANES spectra at Ca L-edge, Sr K-edge and Ba L-edge can convince the 2+ valence state of alkali earth elements and strengthened our XRD results of single phase nature of the samples.

Fig. 5(a) shows the pre/post-edge background subtracted and normalized V K-edge XANES spectra from the V metal foil, VO2, V2O5 and A2V2O7 samples. The first absorption feature between 5465–5475 eV, known as pre-edge peak, originates due to the transitions from 1s core levels to 3d state. Although, it is a dipole-forbidden transition but originates because of the combination of strong 3d–4p mixing and overlap of the metal 3d orbitals with the surrounding O 2p orbitals. Above the edge-energy, a strong peak is present in every spectrum which originates due to the dipole-allowed 1s–4p transitions followed by the other higher energy features from the transitions to higher np states and other multiple scattering contributions or, simply, the EXAFS signals. It is noticeable from the Fig. 5(a) that reference samples show a systematic edge-energy shift towards higher energy with increasing oxidation state of the V ions (i.e., energy position order is V2O5 > VO2 > V). The variation in the energy position is fairly visible from the second derivative spectra of the V K-edge XANES, presented in the Fig. 5(b) and inset therein. The edge-energy variation in the V K-edge spectra is corresponding to the oxidation state changes in the samples.

The change in valence state has been estimated, quantitatively, by plotting the relative energy position of 1s–4p peak to the pre-edge peak as a function of formal oxidation state via deriving an approximate calibration curve and is presented.
in the inset of Fig. 5(a). It is noticeable that the A2V2O7 samples show the sub-pentavalent state of V ions. But the oxidation state of V ions in A2V2O7 samples is comparable (4.62–4.82), regardless of the variation in the alkali-metal element in the compounds. A little less oxidation of V ions, than the expected 5+ oxidation, could be due to the perturbation in the oxygen stoichiometry in the A2V2O7 samples under the given synthesis conditions.

Besides the oxidation state estimation of V ions, the pre-edge peak is prone to the electronic configuration (or empty density of states) and local symmetry of the V ions in various compounds.18,19 The pre-edge intensity will be virtually zero in case of regular octahedral symmetry of the V ions but it enhances for the distorted symmetric compounds and reaches to maximum for the tetrahedral symmetric compounds.19 In the present study, the pre-edge intensity of A2V2O7 compounds is superior to the VO2 or V2O5 samples and is strengthening the more distorted octahedral symmetry or existence of tetrahedral symmetry of V ions in the pyro-vanadates. The Ca2V2O7 compound is exhibiting a lower intensity of pre-edge peak, while compared with Sr2V2O7 and Ba2V2O7, which may indicate lesser transition probabilities from the V 1s core levels to 3d states in this compound compared with other two pyro-

vanadates. Moreover, the pre-edge intensity of V K-edge XANES spectra of A2V2O7 may be subjected to the diverse 3d-4p mixing and overlapping of O 2p with the V 3d orbitals under the influence of alkali metal orbitals. Therefore, to understand the local symmetry of V ions, surrounded by the O ligands in A2V2O7 compounds, it is advantageous to investigate the O K-edge XANES because it gives precise measure of bottom of the conduction band of a compound via probing the unoccupied density of states in t2g and eg orbitals along with the measure of crystal field splitting.

Fig. 6(a) and (b) present the V L-edge and O K-edge XANES spectra of Ca2V2O7, Sr2V2O7, and Ba2V2O7 compounds. As stated above, the d electronic configuration of V d orbitals provides 5+ oxidation state of V ions in A2V2O7 compounds which is similar to the oxidation state of V ions in V2O5 compound (i.e., V5+). Therefore, spectral features and energy position of A2V2O7 compounds were compared with the V L-edge and O K-edge XANES spectrum of the reference V2O5 compound.

According to the dipole selection rules (Δf = ±1), XANES spectra at V L-edge and O K-edge are originated due to the d-projected unoccupied density of states and p-projected density of states of the valence levels, respectively.18–42 There are two distinct peaks in the V L-edge spectra at ~519.5 eV (peak c; known as L3-edge) and 526.5 eV (peak d; known as L2-edge), which are corresponding to the electronic transitions from the V 2p3/2 and V 2p1/2 core levels, respectively, to the unoccupied 3d levels.18,32–40 The V L3 and L2-edges of vanadium compounds obey several spectral features which originate from the crystal field and/or atomic-multiplet effects.18–42 However, the V L2-edge is less important for the analysis because of its broadened nature via the Coster–Kronig Auger decaying process into a 2p3/2 hole.39 Therefore well-resolved V L3-edge features are important to understand the electronic structure assets of the compounds. According to the band-structure calculations of vanadium compounds.
oxides by Maganas et al., the c feature exhibits more of the 3d_2 transition of the V ions whereas the b and a features arise from the 3d_x + 3d_y and 3d_z components, respectively. However, better-resolved spectra of vanadium oxides have also shown another spectral feature between the b and c peaks and it was assigned to the V 2p to V 3d transition. In the present study, we observed two distinct peaks in the V L_3-edge spectra of A2V2O7 compounds (i.e., # and * peaks in the Fig. 6(a)), instead of three features in the V2O5 sample. This indicates that V L_3-edge A2V2O7 compounds is composed of two main transitions between the V 2P_3/2 and 3d orbitals. The # and * peak can be assigned to the transitions between V 2P_3/2 → 3d (x^2, y^2, xz, yz) and V 2P_1/2 → 3d (x^2, y^2, z^2) orbitals. This presumably due to the different crystal field or multiplet effects in the sub-pentavalent A2V2O7 with triclinic crystal structure compared with pentavalent V2O5 with an orthorhombic crystal structure. However, there are no measurable changes in the V L_3-edge spectral features of Ca2V2O7, Sr2V2O7 and Ba2V2O7 samples and are indicating the similar local symmetry (tetrahedral) and oxidation state of V ions as suggested by the V K-edge XANES.

The feature-rich O K-edge spectra are seen in the Fig. 6(b), which are originated from the electronic transitions from O 1s to O 2p hybridized V 3d orbitals and are known to provide valuable information on the crystal structure, local symmetry of metal ions and ligand-field effects. In an octahedral environment, the metal e_g (d_x^2, d_y^2, d_z^2) orbitals are pointed directly towards the O ligands to form strongly bonded σ bonds with the O p_orbital. On the other hand, the metal t_2g (d_{xy}, d_{xz}, d_{yz}) orbitals points between the O ligands and make weak π bonds with the O p_orbital. Consequently, the energy hierarchy of the anti-bonding orbitals is t_2g < e_g and their intensity ratio is 3/2 for an octahedrally coordinated compound. In contrast to the octahedral ligand field, both the symmetries of molecular orbitals and their energy hierarchy are reversed in case of tetrahedrally coordinated compounds. As a result of the tetrahedral interaction of ligands, the hybridized molecular orbitals are characterized by lower lying e_g (π-type) and higher energy-lying t_2g (σ-type) orbitals with their intensity ratio of 2:3.

As stated above, the V ions in V2O5 obey the distorted octahedral symmetry, therefore, the e and f peaks in the O K-edge XANES can be assigned to the t_2g and e_g, respectively. However, the V ions form (VO_4) groups in A2V2O5 compounds with tetrahedral symmetry. Therefore, the energy hierarchy of t_2g and e_g peaks and their intensity ratios are expected to be reversed for the Ca2V2O7, Sr2V2O7 and Ba2V2O7 samples. It can be seen from the Fig. 6(d) that the I(t_2g/e_g) ratio is nearly 3/2 for the V2O5 and is significantly lowered for the A2V2O5 samples. However, the I(t_2g/e_g) ratio for the A2V2O5 samples is not reached up to 2/3 (for the ideal tetrahedral symmetry). This could be due to the availability of octahedral locations to a few of the V ions in A2V2O5 samples along with their tetrahedral occupancy. It is noticeable that, the intensity of the second peak in Fig. 6(c) is higher for the Ca2V2O7 sample (which gives minimum I(t_2g/e_g) ratio) compared with remaining two samples of pyro-vanadates. This is indicating the much-unoccupied density of states were available for the second peak of Ca2V2O7 sample compared with Sr2V2O7 or Ba2V2O7.

In the present study, the Δ_a (crystal field splitting parameter for the pyro-vanadates) is marginally lesser than the reference V2O5 sample (Δ_b). In general, the octahedral ligand field (with six O ions) gives much crystal field splitting (Δ) values while the tetrahedral ligand field (with four O ions) leads to lesser Δ values to a given compound. Therefore, the observed decrease in the Δ_a reinforces the tetrahedral symmetry in the pyro-vanadates as suggested by the V K-edge XANES.

Overall, the V K-edge and O K-edge spectra of pyro-vanadates samples strengthened tetrahedral occupancy of the V ions. However, the Ca2V2O7 sample point a distinct difference in the electronic hybridization of the frontier orbitals of Ca2V2O7 sample compared with Sr/Ba based pyro-vanadates. This could be due to the dissimilar distribution of V ions at tetrahedral/octahedral sites and their hybridization with O 2p orbitals in this compound (compared with other two pyro-vanadate).

c) Photoluminescence study

Fig. 7(a–c) shows the PL spectra of Ca2V2O7, Sr2V2O7 and Ba2V2O7 samples along with their CIE diagrams (g). Fig. 7(d–f) show the...
emission spectra with the Gaussian fittings of emission bands (at energy axis). It is noticeable from the Fig. 8a) that the Ca2V2O7 shows the intense emission bands near to the yellow-orange color (see spot S1 in CIE diagram) and in accordance with the previous reports. The broad emission band centered ~ 530 nm and ~507 nm is observed for the Sr2V2O7 (Fig. 7b) and Ba2V2O7 (Fig. 7c) samples, respectively. This kinds of PL spectra have resulted in the blue-green and blue-indigo emission characteristic in the CIE diagram (Fig. 7g) of Sr2V2O7 (S2) and Ba2V2O7 (S3) samples, respectively.

One of the possibilities of broad-emission spectra could be the CT of an electron from O 2p orbitals to the vacant 3d orbitals of V5+ ions under the tetragonal symmetry of V–O4 groups. In previous reports, it was also assumed that the forbidden luminescence process with the perfect tetragonal symmetry of V–O4 groups would be partially allowed due to the distorted structure as well as the spin-orbit interaction. Additional to that, theoretical calculations were also performed to explain the broad emission spectra of Sr2V2O7:Eu crystals. In such calculations, a 10 × 10 energy matrix was successfully established, based on an effective operator Hamiltonian, including free ion and crystal field interactions. According to the crystal field theory, the pentavalent vanadium (i.e. V5+ ions with 3d⁰ electronic configuration) in the alkali earth metal based pyro-vanadates may not result in any d-d transitions. The d-d electronic transition based emission-characteristics can be achieved from the V5+ ion if it gets one electron from the surrounding oxygen atoms and attains 3d¹ electronic configuration via formation of V4+ ions (i.e., V⁵⁺O₂⁻ → V⁴⁺O₂⁻) (ref. 44). Based on the theoretical calculations, it was conveyed that the broad PL properties of Sr2V2O7:Eu compound originate from four different VO4 groups with C₃v symmetry and not from the f-d transitions of the Eu²⁺ ions. In contrary to this, f-d transition induced broad luminescence has previously reported for the Eu²⁺ and Eu³⁺ dual emission centers. Thus, the mechanistic concept of the broad luminescence spectra from alkali earth metal based pyro-vanadates is still under development.

In the present study, all of the samples have exhibited broad PL properties which can be assigned to the charge transfer (CT) transitions from the O 2p orbitals to the V 3d orbitals within the VO₄ tetrahedra. The molecular orbitals of the V⁵⁺ ions, with the tetrahedral symmetry, obey ¹A₁ ground state and four (¹T₁g, ¹T₂g, ¹T₁u and ¹T₂u) excited states as shown in the Fig. 8(h). Though the all three pyro-vanadates of this study obeyed the triclinic crystal structure, with a space group of P1(2), there are local structural dissimilarities. In the Ca2V2O7, VO₄ dimers approach to each other within the shortest unit cell (compared with the Sr2V2O7 and Ba2V2O7) and the VO₄ pyramids are formed by the manifestation of edge-sharing at adjacent VO₄ tetrahedra. Therefore miscellaneous CT transitions are expected for the Ca2V2O7 compound which, indeed, facilitate yellow-orange emission properties. However, this kind of emission from the A₂V₂O₇ needs electronic structure correlations.

In case of A₂V₂O₇ compounds, the valence band is derived from the fully occupied O 2p orbitals and the conduction band is mainly consist of empty V 3d orbitals. The O K-edge XANES of Sr2V2O7 and Ba2V2O7 samples has shown similar crystal field splitting and nearly the same t₂g/e_g ratio which indicates iso-structured electronic structure properties in these two compound. On the other hand, the t₂g/e_g ratio is significantly decreased for the Ca2V2O7 sample. This indicates that more unoccupied 3d density of states were made available for this compound and could be due to the mixed occupancy of V ions (tetrahedral and/or octahedral). The adverse charge-transfer of an electron from O 2p orbitals to the vacant 3d orbitals of V ions under the distorted tetragonal symmetry may obey the distinct emission properties from the Ca2V2O7. The diverse splitting of V 3d orbitals had previously studied for the Ca2V2O7 (but not seen for the Sr2V2O7 and Ba2V2O7) by the partial density of state (PDOS) calculations and supports our XANES investigation.

The broad emission from the A₂V₂O₇ compounds consist of two distinct emission peaks (higher energy peak (emission 1) and lower energy peak (emission 2)) which are corresponding to the transitions from ³T₂ and ³T₁ to the ground state ¹A₁ (see Fig. 7h). It is evident from the Fig. 7(d–f) that the emission 1 is stronger than the emission 2 and consistent of the previous reports. Moreover, the peak position of emission 1 and emission 2 was red-shifted with decreasing the ionic radii of alkali earth ions. The luminescent colour coordinates of A₂V₂O₇, which were obtained from the CIE chromaticity diagram, were changed from yellow-orange (x = 0.5308, y = 0.4151; for Ca₂V₂O₇), green-blue (x = 0.2445, y = 0.3357; for Sr₂V₂O₇) to blue-indigo (x = 0.2053, y = 0.2666; for Ba₂V₂O₇). The variation in the color coordination in A₂V₂O₇ compounds could be due to the dissimilarities in the electronic/atomic structure as discussed above. In the present study, the CIE coordinates of Sr₂V₂O₇ are quite close to the CIE coordinate of white light (i.e., x = 0.333, y = 0.333). Moreover, advanced emission characteristics may be anticipated by mixing A₂V₂O₇ pyro-vanadates in certain weight/molar ratio and has been

Fig. 8 Emission spectra (λex = 375 nm) of mixed A₂V₂O₇ samples with their equal weight proportions; (a) Ca₂V₂O₇ + Sr₂V₂O₇, (b) Ca₂V₂O₇ + Ba₂V₂O₇, (c) Ca₂V₂O₇ + Sr₂V₂O₇ + Ba₂V₂O₇, (d) Shows the CIE diagram for S1 (Ca₂V₂O₇ + Sr₂V₂O₇), S2 (Ca₂V₂O₇ + Ba₂V₂O₇) and S3 (Ca₂V₂O₇ + Sr₂V₂O₇ + Ba₂V₂O₇) samples.
attempted in the present study by mixing the individual powders in their equal weight percentage.

Fig. 8(a–c) show the PL emission spectra from the mixture of the samples ((a); Ca$_2$V$_2$O$_7$ + Sr$_2$V$_2$O$_7$, (b); Ca$_2$V$_2$O$_7$ + Ba$_2$V$_2$O$_7$, and (c); Ca$_2$V$_2$O$_7$ + Sr$_2$V$_2$O$_7$ + Ba$_2$V$_2$O$_7$) along with their CIE coordinate diagram in Fig. 8(d). It is evidenced from the Fig. 8 that the mixed samples have exhibited much fascinating spectral features and contain the emission bands from constituent compounds. It is noticeable that the intensity of 600 nm–700 nm peaks, which is characteristic yellow-orange emission of Ca$_2$V$_2$O$_7$, is significant in case of the mixture of Ca$_2$V$_2$O$_7$ + Ba$_2$V$_2$O$_7$ sample but has been less dominating in case of other two combinations. This could be due to the superior emission of Ca$_2$V$_2$O$_7$ over the Ba$_2$V$_2$O$_7$ sample. In the Fig. 8(d), samples are named as S1 (for Ca$_2$V$_2$O$_7$ + Sr$_2$V$_2$O$_7$ mixture) S2 (for Ca$_2$V$_2$O$_7$ + Ba$_2$V$_2$O$_7$ mixture) and S3 (for Ca$_2$V$_2$O$_7$ + Sr$_2$V$_2$O$_7$ + Ba$_2$V$_2$O$_7$ mixture). Thus the obtained CIE colour coordinates for S1 (x = 0.3696, y = 0.4617), S2 (0.3561, y = 0.4207) and S3 (0.3462, y = 0.4562) are quite diverse to that of single component pyro-vanadates. The obtained colour coordinates from the mixed compounds are very close to the white-light colour coordinates. Therefore, we may conclude that the vanadium-based compounds of the present study are proficient for the white-light-emitting devices without adding the rare-earth or other dopants.

Conclusions

Single phase pyro-vanadate (Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$, and Ba$_2$V$_2$O$_7$) ceramics were successfully synthesized using a modified chemical precipitation method followed by air annealing of the samples. Ca$^{2+}$, Sr$^{2+}$, and Ba$^{2+}$ ions were confirmed in the Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ samples, respectively, using the XAFS study at the Ca L-edge, Sr K-edge and Ba L-edge. Sub-pentavalent vanadium ions have been confirmed in all three pyro-vanadates of this study by analyzing the V K-edge XANES spectra. O K-edge and V K-edge XANES have confirmed the tetrahedral occupancy in the pyro-vanadate samples. However, the Ca$_2$V$_2$O$_7$ exhibits a smaller t$_{2g}$/e$_g$ peak intensity ratio in the O K-edge which indicates a much-unoccupied density of states in O 2p and V 3d hybridized orbitals under the mixed tetrahe-dr/ocathedral V locations in this sample. Charge-transfer transitions in VO$_4$ tetrahedra have established broad photoluminescence characteristics in pyro-vanadates. Yellow-orange (CEI colour coordinates; x = 0.5308, y = 0.4151), green-blue (CEI colour coordinates; x = 0.2445, y = 0.3357) and blue-indigo (CEI colour coordinates; x = 0.2053, y = 0.2666) emissions have been observed from the Ca$_2$V$_2$O$_7$, Sr$_2$V$_2$O$_7$ and Ba$_2$V$_2$O$_7$ samples, respectively. The emitted colour coordinates could be tailored by mixing of the compounds and has resulted in advanced photoluminescence properties which are comprehensively suitable for the rare-earth element free white-light-emitting devices.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

18. R. Yu, M. Yuan, T. Li, Q. Tu and J. Wang, RSC Adv., 2016, 6, 90711.