# **RSC Advances**



**PAPER** 

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2017, 7, 8131

# Positron annihilation studies and complementary experimental characterization of $xAg_2O-(1-x)(0.3CdO-0.7MoO_3)$ metal oxide glass nanocomposites†

Ranadip Kundu,‡\*a Sanjib Bhattacharya,b Debasish Roya and P. M. G. Nambissanc

Metal oxide nanocomposites of the composition  $xAg_2O-(1-x)(0.3CdO-0.7MoO_3)$  were prepared by a melt-quenching method and were characterized by different experimental techniques like X-ray diffraction, high resolution transmission electron microscopy and optical absorption spectroscopy. X-ray diffraction showed sharp diffraction peaks indicating large crystallites but transmission electron microscope images also showed crystallites of nanometer dimensions in appreciable concentrations, which confirmed the nanocomposite structure of the samples. Although the lattice constants did not show significant changes with the increase in concentration (x) of Aq<sub>2</sub>O, there is considerable relaxation of the growth-induced strain above x = 0.2. Interestingly this is also the concentration above which the optical band gap energy showed a mild decrease. One salient feature of this study is the use of positron annihilation spectroscopy for identifying and monitoring the structural defects such as vacancies and vacancy clusters as well as the free volume cavities during the change in concentration of Aq<sub>2</sub>O. Positron lifetime measurements indicated trapping of positrons initially in the interfacial defects within the  $0.3CdO-0.7MoO_3$  nanocrystalline ensemble and then in the free volume defects within the amorphous Ag<sub>2</sub>O matrix. At higher Ag<sub>2</sub>O concentrations, positron trapping appeared to take place within the Cd<sup>2+</sup>-monovacancies in CdO and in the divacancies of neighbouring cationic and oxygen monovacancies in the  $\alpha$ -MoO<sub>3</sub> and CdMoO<sub>4</sub> nanocrystallites. At x=0.1-0.2, the effective positron trapping centres are translocated to the tetrahedral Mo<sup>6+</sup>-monovacancies instead of the Cd<sup>2+</sup>monovacancies. The results of coincidence Doppler broadening spectroscopic measurements, which map the electron momentum distribution and its variations, indicated increasing trapping of positrons with increasing concentration of Ag<sub>2</sub>O, which again is attributed to the trapping sites in the increasing number of nanocrystallites being formed

Received 23rd September 2016 Accepted 13th January 2017

DOI: 10.1039/c6ra23680d

www.rsc.org/advances

### Introduction

Metal oxide based glass nanocomposites and other similar materials have gained popular importance in recent years as the discovery of nanocrystalline properties of materials have opened up new avenues of research. Of particular interest is the tunability of such materials with the introduction of substituted

elements that can enhance or modify their physical and other properties to the extent of adaptability in technological applications. Glassy systems and their nanocomposites are of special interest and focus in this context as they cater to a wide range of applications such as solid state batteries and sensors.1,2 Understanding their properties and the factors affecting their variations under different thermodynamic and experimental conditions is therefore really important. In this context, we report here the results of a very recent study on the electronic and stoichiometry-related properties of a glass nanocomposite consisting of Ag<sub>2</sub>O and 0.3CdO-0.7MoO<sub>3</sub> in different proportions with a general nomenclature  $xAg_2O-(1 - x)(0.3CdO 0.7\text{MoO}_3$ ). The variational parameter x had values starting from 0.0 to 0.9. Altogether seven samples have been studied and the modifications of the structure and properties have been investigated.

The findings from conventional experimental techniques like Fourier transform infra-red (FTIR) and UV-Vis absorption

<sup>&</sup>quot;Department of Mechanical Engineering, Jadavpur University, Jadavpur, Kolkata 700032, India. E-mail: debasish\_kr@yahoo.co.in

<sup>&</sup>lt;sup>b</sup>Department of Engineering Sciences and Humanities, Siliguri Institute of Technology, Darjeeling 734009, West Bengal, India. E-mail: sanjib\_ssp@yahoo.co.in

<sup>&</sup>lt;sup>c</sup>Applied Nuclear Physics Division, Saha Institute of Nuclear Physics, Kolkata 700064, India. E-mail: pmg.nambissan@saha.ac.in

<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: 10.1039/c6ra23680d

<sup>‡</sup> Present address (RK): Department of Electrical Engineering, University of Engineering and Management, Kolkata-700156, West Bengal, E-mail: ranadip.89@gmail.com; Fax: +91-33-23578302; Tel: +91-8902390462.

spectroscopies are also often complemented and supported by characterization tools like X-ray diffraction (XRD) and high resolution transmission electron microscopy (HRTEM).

However, there are bigger challenges like the deviation of measured electrical conductivity from the theoretically expected ones and the modifications of thermal properties which needed deeper understandings and evaluation. Of special emphasis are the defects and defects-related properties and processes that may as well undergo evolution with the variations of the concentrations of the constituents in the composition of the nanocomposites. Positron annihilation spectroscopy is a versatile non-destructive defect spectroscopic probe that has been increasingly applied in the investigation of the defects in nanomaterials and nanocomposites.3-5 Positron lifetime (PL) measurements as well as coincidence Doppler broadening spectroscopy (CDBS) were performed on the seven samples and the results are discussed.

# Experimental details

**RSC Advances** 

The samples were prepared through a chemical mixing process using reagent grade chemicals AgNO<sub>3</sub>, CdO and α-MoO<sub>3</sub>. Depending upon the molar fraction x of the Ag<sub>2</sub>O part in the stoichiometric composition, calculated amounts of the reagents have been accurately weighed and taken in an alumina crucible and thoroughly mixed and preheated. (See ESI I.†) The mixtures were then melted in an electric furnace in the temperature range from 900 °C to 950 °C depending upon the composition. The melts were equilibrated for 30 minutes and then quenched between two aluminum plates. In this manner, glassnanocomposites of thickness  $\sim 1$  mm have been obtained for x = 0.0 (without AgNO<sub>3</sub>)-0.9. For experiments in the powder form of the samples, the plates so formed are gently crushed in an agate mortar and filtered through a 100  $\mu m$  sized mesh.

X-ray diffraction (XRD) patterns of the samples were recorded using  $CuK_{\alpha}$  radiation (wavelength 1.54 Å) in a rotating anode Rigaku TTRAX-III X-ray diffractometer. Scanning is done from  $2\theta = 10^{\circ}$  to  $80^{\circ}$  through steps of  $0.02^{\circ}$  with a scanning speed of 4° per minute. The presence and distribution of the nanocrystallites have been confirmed from the HRTEM images taken using 300 keV electrons (FEI, Tecnai, F30-ST). UV-Vis absorption studies were made in the reflection mode using PerkinElmer Lamda-750 spectrophotometer. The colloidal suspensions using ethanol as solvent for the absorption studies were prepared with the help of an ultrasonic homogenizer (Takashi SK-500F).

For positron annihilation studies, the fine powdered sample is taken in a glass tube and a radioactive isotope decaying by  $\beta^+$ emission is kept embedded in it at the geometrical centre of the column. The source in fact is a 22NaCl deposition on a thin  $(\sim 2 \text{ mg cm}^{-2})$  Ni foil and covered by the extended portion of the foil folded along a line away from the source-deposited area. Prior to the source deposition, the Ni foil had been annealed at 1050 °C for 2 hours in good vacuum ( $\sim 10^{-6}$  mbar). The powdered sample is ensured to cover the source from all sides in thicknesses sufficient to capture and annihilate all the positrons within it and none of them reached the glass walls.

Further, the glass tube is continuously evacuated to remove the air and absorbed gases from the powder column and to allow the particles to settle down under their own weight. In this way, the sample and the source assembly is made compact and maintained at dry and moisture-free conditions. The positron emission and annihilation gamma rays are captured using nuclear radiation detectors and processed by standard gammagamma coincidence circuits, comprising of nuclear electronic modules.4 Special mentions are to be made of the detectors used for the detection process. For positron lifetime measurements, BaF2 scintillators coupled with fast XP2020Q photomultiplier tubes were used. The slow-fast gamma-gamma coincidence spectrometer which incorporated these detectors and the associated nuclear electronic modules had a time resolution of 180 ps when measured with the nuclear gamma rays from a 60Co radioactive isotope kept under the 22Na experimental energy window settings. For CDBS studies, two high pure germanium (HPGe) detectors with energy resolutions  $\sim$  1.27 keV at the annihilation gamma ray energy 511 keV are used. The data were collected with sufficiently high statistics  $(\sim 10^6 \text{ and } 10^7 \text{ counts or events under each PL and CDB spectra})$ respectively) and the electronic stability had been extremely good. The positron lifetime spectra were analyzed using the program PALSfit.6 The software LAMPS developed by Pelletron Division, Tata Institute of Fundamental Research, Mumbai<sup>7</sup> was used in the CDBS data acquisition, storage and analysis.

### 3. Results and discussion

### 3.1. X-ray diffraction

The XRD patterns of the synthesized nanocomposite samples are shown in Fig. 1(a). All the spectra displayed sharp welldefined peaks and the identification of the different diffraction peaks had been carried out by a careful comparison with the available literature data of CdO (ICDD PDF: 75-0594) and  $\alpha$ -MoO<sub>3</sub> (ICDD PDF: 05-0508). As reported by Usharani et al.,8 Stoyanova et al.9 and Alsaif et al.,10 the sharp peaks at  $2\theta =$ 12.85°, 27.38° and 39.05° emerge due to the diffraction from the (0 2 0), (0 4 0) and (0 6 0) planes of the large number of crystallites preferentially oriented in these planes in a lamellar configuration. Whereas, the peak of the maximum intensity corresponding to the (1 1 1) plane of CdO appears at  $2\theta = 32.08^{\circ}$ and that of the (1 1 0) plane of α-MoO<sub>3</sub> makes its presence visible at  $2\theta=23.41^{\circ}$ . There are however a number of additional diffraction peaks which lay partially hidden due to the prominent intensities of the (0 2 0), (0 4 0) and (0 6 0) peaks and they are shown magnified in Fig. 1(b). These peaks are observed at  $2\theta$  $= 29.27^{\circ}$ , 35.04°, 48.14° and 51.80° and identified as the (1 1 2), (2 0 0), (1 2 3) and (2 2 0) peaks of cadmium molybdate (CdMoO<sub>4</sub>) (ICSD #084455), having a tetragonal scheelite-type structure with space group  $I4_1/a$  and composed mainly of CdO<sub>8</sub> octahedra and MoO<sub>4</sub> tetrahedra. 11,12 The procedure of synthesis adopted in this work seemed to have given favourable thermodynamic conditions for the formation of this phase, albeit unexpectedly. A still deeper scrutiny of the diffraction patterns also revealed very small peaks at  $2\theta = 33.82^{\circ}$ ,  $55.35^{\circ}$ , 65.03° and 77.65°, which were identified as due to the (1 1 1), (2

Paper

x Ag<sub>2</sub>O-(1-x)(0.3CdO-0.7MoO<sub>3</sub>)

(a) x=0.9

x=0.7

x=0.5

x=0.1

x=0.1

2θ (Degree)

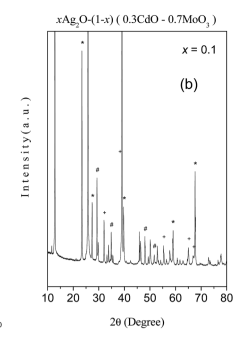


Fig. 1 (a) The XRD patterns of the samples. (b) Gives the XRD pattern of one of the samples with a higher magnification in the y-axis. The symbols +, \* and # respectively indicate the peaks from the three nanocrystalline phases CdO,  $\alpha$ -MoO $_3$  and CdMoO $_4$ . The unmarked peaks mainly belong to Ag $_2$ O and a few of them to Ag.

2 0) and (3 1 1) planes of  $Ag_2O$  (ICDD PDF: 00-065-6811) and (3 1 1) plane of Ag (ICDD PDF: 00-004-0783). Since the intensities of these peaks are found extremely low, they are not considered for further discussion.

The results of XRD analysis did not indicate considerable widths for the peaks and it demonstrated the coarse dimensions ( $d_c$ ) of the large crystallites. The Scherrer equation<sup>13</sup>

$$d_{\rm c} = \frac{0.9\lambda}{\beta_{hkl}\cos\theta} \tag{1}$$

where  $\lambda$  is the wavelength of the  $\text{CuK}_{\alpha}$  radiation was used to calculate the sizes of the crystallites of the different solid phases formed.  $\beta_{hkl}$  is the full width at half maximum of the respective peaks after the correction for broadening due to finite instrumental resolution is incorporated. The average sizes of the crystallites so estimated were large ( $\sim 300-500$  nm) and raised ambiguities related to the nanocomposite nature of the samples. Still the sharp and well-defined peaks confirmed the purity of the phases formed and the good crystallinity achieved.

The lattice constants of the crystallites of CdO,  $\alpha$ -MoO $_3$  and CdMoO $_4$  have also been calculated by considering the respective structure as reported in literature and the identification of the positions of the X-ray diffraction peaks from Fig. 1. The results are summarized in Table 1. No significant variations are observed in the lattice constants. This is interesting especially in the context of a number of reports on the size-dependent lattice parameter changes in a number of metal oxide nanocrystalline systems. The co-existence of three chemically stable metal oxides with homogenous distribution and forming

Table 1 The lattice constants of the nanocrystallites of CdO,  $\alpha$ -MoO $_3$  and CdMoO $_4$  estimated from the XRD patterns shown in Fig. 1

	~ (Å)	L (Å)	- (Å)
Concentration of $Ag_2O(x)$	a (Å)	b (Å)	c(A)

Crystallite: CdO; most intense peak at  $2\theta=33.20^\circ$ ;  $(h\ k\ l)=(1\ 1\ 1)$ ; structure: fcc; ICDD PDF: 75-0594; reported lattice constant a=4.686 Å

0.0	4.668
0.1	4.667
0.2	4.670
0.3	4.672
0.5	4.665
0.7	4.670
0.9	4.670

Crystallite:  $\alpha$ -MoO<sub>3</sub>; most intense peak at  $2\theta = 25.79^{\circ}$ ; ( $h \ k \ l$ ) = (0 4 0), structure: distorted orthorhombic; ICDD PDF: 05-0508; reported lattice constants a = 3.963 Å, b = 13.855 Å, c = 3.696 Å

0.0	3.950	13.804	3.680
0.1	3.946	13.800	3.686
0.2	3.950	13.812	3.685
0.3	3.953	13.818	3.692
0.5	3.945	13.794	3.685
0.7	3.947	13.805	3.686
0.9	3.948	13.806	3.687

Crystallite: CdMoO<sub>4</sub>; most intense peak at  $2\theta=29.30^\circ$ ; (h k l) = (1 1 2); structure: regular orthorhombic; ICSD #084455; reported lattice constants a=5.155 Å, c=11.194 Å

COLIDCCITED OF	0.100 11, 0	11117111	
0.0		5.139	11.150
0.1		5.138	11.146
0.2		5.144	11.146
0.3		5.146	11.160
0.5		5.137	11.146
0.7		5.139	11.159
0.9		5.141	11.156

**RSC Advances** 

a glass-like matrix could have resulted into the mutual annulment of the changes in this case.

We also calculated the lattice strain associated with the formation of the nanocrystallites within the glass matrix. The contribution of the strain to the broadening of the peaks in the XRD patterns of the samples due to crystal imperfections and dislocations is given by

$$\varepsilon = \frac{\beta_{hkl}}{4 \tan \theta} \tag{2}$$

The basic difference between the crystallite size and the induced lattice strain is the nature of their dependency on the diffraction angle  $2\theta$ . The crystallite sizes follow  $1/\cos\theta$  variation whereas the strain varies as  $1/4 \tan \theta$ . On the basis of the independent contribution of each other to the peak broadening, the strain is estimated through the Williamson-Hall (W-H) method.15 The total broadening of the peak is written as the sum of contributions of both the effects as

$$\beta_{hkl} = \frac{0.9\lambda}{d_c \cos \theta} + 4\varepsilon \tan \theta \tag{3}$$

Rearranging the terms, we get

$$\beta_{hkl}\cos\theta = \frac{0.9\lambda}{d_c} + 4\varepsilon\sin\theta \tag{4}$$

Eqn (4) represents the uniform deformation model where the strain induced by the nanocrystallites is assumed to be uniform. From the linear plot of  $\beta_{hkl}\cos\theta$  versus  $4\sin\theta$ , the strain is obtained from the slope of the line. (The y-intercept gives the sizes of the nanocrystals, as given by eqn (1).) A plot of the values of the strain thus estimated versus the concentration (x) of Ag<sub>2</sub>O of the different samples is given in Fig. 2. Although the magnitudes of the strain thus estimated are extremely small, the striking observation of an abrupt discontinuity of the strain at x = 0.2 to 0.3 is worth noting.

### 3.2. TEM images

The X-ray diffraction peaks had been rather sharp giving doubts to the formation of very large crystallites against the nanocomposite nature that was expected. The crystallite sizes estimated from the Scherrer formula also gave the indications that the crystallites are rather large in sizes and the nanocomposition is not to be expected. For disseminating the ambiguity in this regard, we have also examined the samples through high resolution transmission electron microscopy (HRTEM). The TEM images shown in Fig. 3(a) to (c) showed rodlike structures with average diameters showing a gradual increase from 300 nm to 600 nm in samples of increasing Ag<sub>2</sub>O content (x) from 0.1 to 0.9. The diameters of these rods are indicated in the images. The corresponding selected area diffraction (SAD) patterns are shown in Fig. 3(d)-(f) respectively from regions marked by the red circles. The SAD patterns confirm the single crystalline nature of the rod-like structures. The lattice images shown in Fig. 3(g)-(i) further confirm the

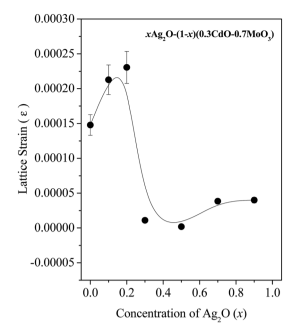


Fig. 2 The crystal lattice strain versus concentration (x) of Ag<sub>2</sub>O used in the samples.

formation of the different solid phases discussed in the previous section.

The TEM images also demonstrated that, in addition to the large rod-like single crystals, there are innumerable nanocrystallites too (Fig. 4), the presence of which in fact should have given rise to finite line broadening for the XRD patterns, had the bulk crystallites been absent. These smaller nanocrystallites are sometimes attached to bigger crystals as, for example, reflected in Fig. 3(e) showing polycrystalline-like diffraction pattern. Further, these nanocrystallites are also single crystals and one can observe lattice plane bending etc. in the Fourier-filtered images shown in Fig. 5(a)-(c). They are also found to have defects such as stacking faults and misfit dislocations associated with them.

### **UV-Vis absorption results**

Although different solid phases are found to be formed within the nanocomposites, they also exhibited a cumulative optical behavior which could be examined and quantified in terms of the band gap energies estimated using Tauc's plots.16 The UVvisible absorption spectra of the different samples are shown in Fig. 6. The tangents to the absorption edges were linearly extrapolated to the x-axis in order to calculate the band gap energies from the relation  $\alpha h \nu = A(h \nu - E_g)^n$  where A is a constant and  $h\nu$  is the photon energy. The value of n depends on the nature of transition.  $^{16}$  In this case, n is chosen as 1/2since both CdO and MoO3 are direct band gap semiconductors and thus the optical band gap energies of the samples are obtained by extrapolating the linear portion of the  $(\alpha h \nu)^2$  vs.  $h \nu$  curve to 0 on the y-axis. The band gap energies thus obtained for the present samples are shown in Fig. 7(a). The initial increase followed by a decrease with increase in

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Paper

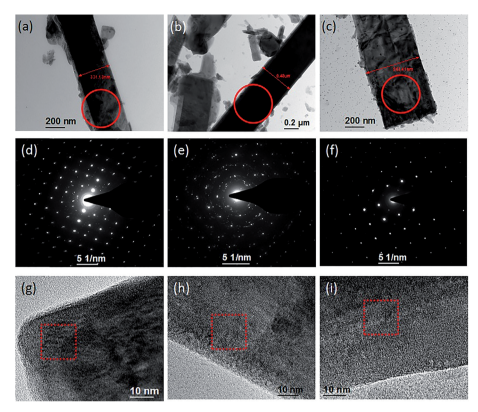


Fig. 3 (a)–(c) TEM images of the three samples with x = 0.1, 0.5 and 0.9. (d)–(f) The SAD patterns. (g)–(i) HRTEM images of the samples showing the lattice fringes.

concentration x of  $Ag_2O$  may more have to do with the actual sizes of the nanocrystallites than any compositional dependence.

With this prelusion, the band gap was plotted as a function of the combined average of the sizes of the nanocrystallites of the three different phases CdO, α-MoO<sub>3</sub> and CdMoO<sub>4</sub> and the results are shown in Fig. 7(b). There is a gradual decrease of the band gap with the increase of the crystallite size. It is not possible here to attribute the increase of band gap with the decreasing crystallite size to quantum confinement behavior (weak, intermediate or strong) since the realm of crystallite sizes for the observation of such effects normally is much below, say  $\sim$ 10 nm or below. For quantum confinement effects to manifest as an increase of the band gap, the nanocrystallite sizes need to

be less than the bulk exciton Bohr radius of the material and the large crystallite sizes obtained in the present study will rule out such a possibility. For example, the exciton Bohr radius for CdO

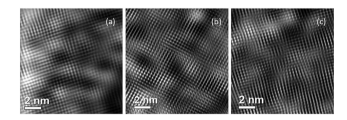


Fig. 5 (a)-(c) Fourier-filtered images of the regions marked in Fig. 4(g)-(i).

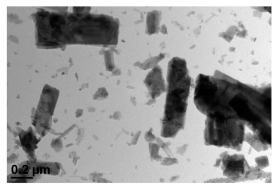


Fig. 4 TEM image of sample with x = 0.5.

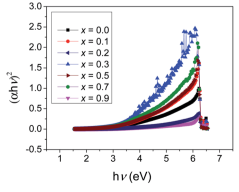


Fig. 6 The UV-Vis absorption spectra of the different samples.

RSC Advances Paper

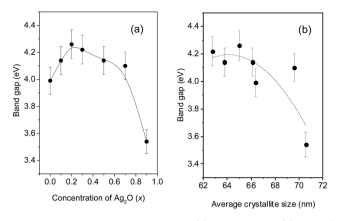


Fig. 7 The calculated band gap *versus* (a) concentration (x) of  $Ag_2O$  used in the samples and (b) cumulative average sizes of the crystallites of the three phases CdO,  $\alpha$ -MoO $_3$  and CdMoO $_4$  obtained in the different samples.

has been reported as close to 2.25 nm (ref. 17) and the values for  $\alpha\textsc{-MoO}_3$  or CdMoO $_4$ , although not readily available in literature, are likely to be of the same order of magnitude. The decreasing band gap within the small range of variation of the crystallite sizes hence can be treated as an "impurity effect" in which additional energy levels are created within the band gap and weak electronic excitations are activated with photons of the corresponding wavelength getting absorbed.

### 3.4. Positron lifetime studies

The peak-normalized positron lifetime spectra of some of the samples are shown in Fig. 8. Notwithstanding the very small variations in the slopes of the curves, the slow decay nature indicates the presence of extended defects of the vacancy-type in the nanocomposites. In solid systems of nanocrystalline composition, the thermal diffusion of the positrons to the outer surfaces of the crystallites also will greatly influence the annihilation characteristics. In the present case, this may not be a serious issue since the sizes of the nanocrystallites are all well above the thermal diffusion lengths of positrons in oxide materials.<sup>18</sup>

The positron lifetime spectra had been analyzed using the PALSfit program to get the different positron lifetimes and their relative intensities. The program subtracts a linear background

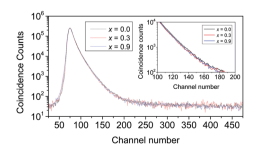


Fig. 8 Peak-normalized positron lifetime spectra of some of the samples. The inset shows a zoomed-in portion of the decaying exponential part for the sake of clarity.

and deconvolutes the resolution function (fitted to a Gaussian) before the necessary source correction (23.75%) is carried out. The source component included the contributions of positrons annihilating within the supporting Ni foil (0.100 ns; 69.83%) and the  $^{22}$ NaCl source material (0.385 ns; 29.62%) and those getting backscattered (1.960 ns; 0.55%). These values were obtained from the analysis of the spectra earlier acquired using the same source with high pure (99.999%) and well-annealed Al single crystals. The best fit (with a reduced chi-square = 1.05  $\pm$  0.15) for the multi exponential decay of the spectra of the glass-nanocomposite samples was obtained for three lifetime components. The longest component  $\tau_3$  was of relatively small intensities (0.1–6.0%) but its presence could not be ignored for want of a reasonably good fit in the analysis. The results of the analysis are illustrated in Fig. 9(a) and (b).

The longest component  $\tau_3$  is attributed to the formation and eventual annihilation by "pick-off" process of *ortho*positronium (*o*-Ps) atoms from the free volume defects of the amorphous Ag<sub>2</sub>O matrix of the nanocomposites. The presence of this component is of special significance because it directly provides a method to calculate the size and concentration of the free volume defects. <sup>19</sup> Also, such amorphous material systems are normally abundant with free volume cavities, whose sizes could vary over a few angstroms.

The question about the origin of the third lifetime  $\tau_3$  is to be answered first. The well-known Tao–Eldrup equation<sup>20,21</sup>

$$\tau_3 = 0.5 \left[ 1 - \frac{R}{R_0} + \frac{1}{2\pi} \sin\left(\frac{2\pi R}{R_0}\right) \right]^{-1} \tag{5}$$

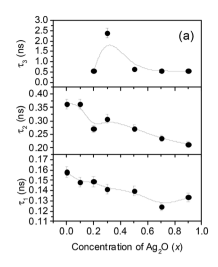
is used to calculate the radius R of the free volume defects and it puts a lower limit of 0.5 ns on  $\tau_3$ . (In eqn (5) above,  $R_0 = R + \Delta R$ where  $\Delta R$  is the empirical electron layer thickness set equal to 1.66 Å. The lifetime  $\tau_3$  is to be expressed in nanoseconds.) In the sample with x = 0.0, due to the absence of Ag<sub>2</sub>O, no third lifetime component had been obtained and the same happened in the next sample with x = 0.1 in which the concentration of Ag<sub>2</sub>O is very small. With increasing concentration to x = 0.2, free volume defects of radius R = 0.55 Å corresponding to  $\tau_3 = 0.550$ ns are being observed and it rose to 3.18 Å in the sample with x= 0.3 where  $\tau_3$  is found as high as 2.382 ns. Subsequently,  $\tau_3$ again fell back to 0.550 ns but the corresponding intensity  $I_3$ gradually rose to 6.0% in the sample of x = 0.9 (Fig. 9(b)). Incidentally, R = 0.55 Å is the lower limit of the free volume cavity size in which o-Ps formation is possible, as reported by Uedono et al.22

The free volume fraction  $(f_V)$  is another quantity which is normally estimated for identical systems by using the relation

$$f_{\rm V} = CV_{\rm f}I_{\rm 3} \tag{6}$$

and its variation with the concentration of  $Ag_2O$  in the samples is shown in Fig. 10. In the above relation,  $C = 0.0018 \text{ Å}^{-3}$  is a constant and  $V_f = (4/3)\pi R^3$  is the geometrical volume (in ų) of the free volume defect considered to be spherical in shape. The significance of this quantity is that it helps to correlate the cumulative free volume fraction with the nature of role of the amorphous matrix aiding or inhibiting the physical properties

Paper



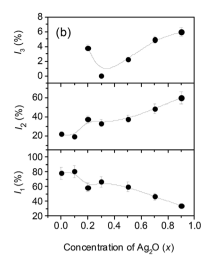


Fig. 9 (a) The positron lifetimes and (b) the relative intensities versus the concentration of Aq<sub>2</sub>O in the samples.

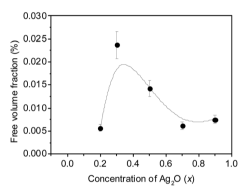


Fig. 10 The free volume fraction versus the  ${\rm Ag_2O}$  concentration of the samples.

associated with them. Thus we can identify that, for the concentration x = 0.2 of Ag<sub>2</sub>O, the degree of amorphousness is rather small and the defects which trap positrons and facilitate the formation of o-Ps atoms are the pores of the amorphous matrix. At x = 0.3, the situation becomes ripe for the formation of fully grown free volume cavities as the bondage between the Ag<sub>2</sub>O matrix and the 0.3CdO-0.7MoO<sub>3</sub> nanocrystalline ensemble is still weak. However, the situation changes again when the concentration of Ag<sub>2</sub>O is increased beyond this threshold when it starts acting like a catalyst to intensify the glass formation by bonding the former ( $\alpha$ -MoO<sub>3</sub>, in this case) with the modifier (CdO) in increasing number. As a result, the free volume fraction is drastically reduced (Fig. 10). However, since the amount of Ag<sub>2</sub>O has increased, there are more number of pores of the smallest sizes and the intensity  $I_3$  accordingly increases to 6%. Thus it could be concluded from the above observations that, while the formation of free volume defects is very well a characteristic of the increasing amorphous Ag<sub>2</sub>O matrix, the formation of o-Ps is strongly inhibited by the presence of the nanocrystallites.

The intermediate positron lifetime component  $\tau_2$  is a parameter of interest in the understanding of the defects

present within the nanocomposite system under investigation. Its value 0.363 ns in the initial two samples (i.e., x = 0.0 and 0.1) is attributable only to positron trapping in very large defects like interfacial vacancy clusters within the 0.3CdO-0.7MoO<sub>3</sub> nanocrystalline ensemble. At these two extremely low concentrations of Ag<sub>2</sub>O, strong bondage between the former and the modifier is absent. The subsequent decrease of  $\tau_2$  by about 0.093 ns is significant as it signals to the formation of the nanocomposite glass due to the development of strong bonding between the α-MoO<sub>3</sub> former and the CdO modifier mediated by the presence of Ag<sub>2</sub>O which acts as a catalyst. As the bonding intensifies, the interfaces that acted as the potential positron trapping sites disappear and positrons instead get trapped into the relatively shallow defects like the negatively charged cationic monovacancies within the nanocrystallites being formed. Thus the decrease of  $\tau_2$  at x = 0.1–0.2 signifies a translocation of the positron trapping sites from the interfaces to the isolated cationic monovacancy-type defects. As already stated, we could identify from the X-ray diffraction patterns that there are crystallites of CdO, α-MoO<sub>3</sub> and CdMoO<sub>4</sub> but no reported values of positron lifetimes in these composite phases are available so far. On the other hand, the obtained  $\tau_2$  values agree with the values available in literature for the cationic vacancies of some of the elements constituting the above mentioned nanocrystallites. For example, Seeger has reported the positron lifetime in neutral cadmium monovacancies ( $V_{Cd}$ ) as 0.256 ns (ref. 23) whereas, for the negatively charged Cd2+-monovacancies  $(V_{\rm Cd}^{2-})$  in semiconductors like CdSe and CdTe, the reported positron lifetimes varied as widely as from 0.288 ns to 0.325 ns.3 On the other hand, the positron lifetime in neutral monovacancies in silver is 0.208 ns (ref. 24) while a paper published by Wegner<sup>25</sup> suggested 0.180-0.190 ns as the positron lifetime in a silver monovacancy associated with an oxygen ion. Identical extrapolation of these values would then suggest a positron lifetime around 0.235 ns for a Cd<sup>2+</sup>-monovacancy in CdO. It should be mentioned here that positrons are insensitive to the oxygen vacancies  $(V_{\rm O}^{2+})$  because of the positive charge

**RSC Advances** Paper

associated with the latter and they normally get trapped only in neutral or negatively charged vacancies. The values reported for the positron lifetimes in other cationic defects such as Mo or Ag monovacancies are much less, 170 ps and 208 ps respectively. Interestingly, this reference by Campillo Robles and coworkers<sup>26</sup> has also quoted the positron lifetime in neutral Cdmonovacancies as 252 ps (experimental). We therefore attribute the decreasing  $\tau_2$  values in the rest of the samples (i.e., for which x > 0.2) as due to positron trapping in Cd<sup>2+</sup>-monovacancies as well as in  $Mo^{6+}$ -monovacancies. At x = 0.9, the value of  $\tau_2$  is 0.212 ns, which is close to that reported for the monovacancies in Ag<sub>2</sub>O.<sup>24</sup> This is the sample with the highest concentration of Ag<sub>2</sub>O and it shows the presence of neutral crystalline monovacancies. Note that the X-ray diffraction data showed well-defined narrow peaks of the Ag<sub>2</sub>O phase with feeble intensities instead of a broad one, indicating small concentrations of crystalline phases within the Ag<sub>2</sub>O matrix.

It is worth pointing out that metal oxide nanocrystallites are not in general free of vacancy-type defects. These defects are inherent characteristics of any metallic oxide  $(M_m O_n)$  system. Maintenance of an exact stoichiometry of the metal and oxygen is seldom observed and they may be actually described as  $\mathbf{M}_{m-a}\mathbf{O}_{n-b}$  where the deviations a and b vary over a few tens to hundreds parts per million (ppm). Positrons are known to undergo saturation trapping if only the defect concentrations exceed 10<sup>20</sup> cm<sup>-3</sup> which, for example, is about 10<sup>3</sup> ppm in CdO.<sup>3</sup> The unoccupied lattice sites constitute a defect concentration less than this limit and such vacancies are also responsible for the trapping and annihilation of positrons. These vacancies are perhaps inherently built up in the nanocrystallites in order to give them a certain degree of stability and such nonstoichiometric vacancies have been identified as major defect sites in several oxide nanocrystalline systems that have been investigated by a large number of research groups. 27-29

It should be additionally noted that, in such ionic compounds, the vacancy type defects can have electrical charges associated with them. A metallic ion is positive and hence the vacancy generated by its absence will be negatively charged. Positrons will be effectively and efficiently trapped by them. Contrary to this, an oxygen vacancy will be having a positive charge and hence the positrons, which are positively charged particles, cannot be trapped by the oxygen vacancies. Then there is the possibility of a divacancy. In this case, the divacancy being created by the absence of a metallic ion and its immediate neighboring oxygen ion, the two opposite charges nullify each other and the divacancy becomes electrically neutral. Positrons can be still trapped and one gets a characteristic positron lifetime corresponding to it. Hence the second positron lifetime component  $\tau_2$  can be safely attributed to vacancy type defects within the nanocrystallites.

The shortest lifetime  $\tau_1$  is a result of positron annihilation in bulk material and reduced by the free-state residence time of the trapped positrons.<sup>30,31</sup> In the case of the two samples of x =0.0 and 0.1 where the positron lifetime spectra yielded only two components and in the spectra of the sample with x = 0.3, in which the intensity  $I_3$  was just 0.1%, we applied the two-state trapping model<sup>31</sup> to calculate the positron lifetime within

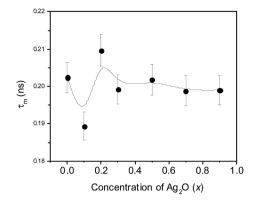


Fig. 11 The mean positron lifetimes versus concentration (x) of Ag<sub>2</sub>O of the samples

a possible "bulk" glass crystallite and it emerged in the range 0.170-0.180 ns. In the rest of the samples, the model did not exactly reproduce the bulk positron lifetime and hence the original scheme of analysis with a distinct  $\tau_3$  and  $I_3$  is retained. Still, in all the cases, there is a contribution of the trapped positrons, whose lifetimes are defect-specific and which will vary according to their sizes and locations. In order to make this discussion complete, we are presenting in Fig. 11 the variation of the mean positron lifetime  $\tau_{\rm m}$  defined as

$$\tau_{\rm m} = (\tau_1 I_1 + \tau_2 I_2 + \tau_3 I_3) / (I_1 + I_2 + I_3) \tag{7}$$

The features exhibited by this parameter is supportive of the previous interpretations with respect to positron trapping being translocated to the vacancy-type defects within the nanocrystallites rather than at the interfaces at higher Ag<sub>2</sub>O concentrations, especially at x > 0.3.

### Coincidence Doppler broadening spectroscopic studies

The electron momentum distributional variations reflected in the events of positrons annihilating with electrons localized at different atomic orbits are studied using the coincidence Doppler broadening measurement procedure described by Asoka-Kumar et al. 32 and widely used by other researchers. 33 The ratio curves generated for the different samples with respect to pure and well-annealed Al single crystals are presented in Fig. 12. The characteristic peak at the longitudinal component of the electron momentum  $(p_{\rm L})$  at  $\sim 10 \times 10^{-3} m_0 c$  comes from positron annihilation by the 2p electrons of oxygen ions. This means the annihilation events contributing to this peak are due to those positrons which are getting trapped at the vacancies created by the absence of the metallic ions at some of the lattice sites within the crystallites. To sense the changes in the cationic distribution, the positrons are to be localized at the oxygen vacancies, which however is not possible due to the Coulomb repulsion faced by positrons. The position of the peak did not show any shift with changes from sample to sample (Fig. 12), indicating that the elemental surroundings around the positron trapping defects remained oxygen throughout. However, the Paper

1.625 1.500 1.375 1.250 1.250 1.250 1.000 0.875 0.750 0.0 2.5 5.0 7.5 10.0 12.5 15.0 17.5 20.0 22.5  $\rho_L(10^{-3} \, \text{m}_0\text{c})$ 

Fig. 12 The CDBS ratio curves of the different samples with respect to single crystalline Al. The curve for pure  $Aq_2O$  (x = 1.0) is also shown.

amplitudes (heights) of the peak shown in Fig. 13 are found to vary in a manner that exhibited a strong correlation with that of the positron lifetimes and intensities (Fig. 9(a) and (b)). Increased annihilation with the electrons of oxygen when the concentration (x) of Ag<sub>2</sub>O increased systematically is the overall trend; however, a characteristic discontinuity at x=0.2–0.3 is remarkable and seems to indicate a sudden change of the sites of positron trapping to one with reduced number of coordinated O<sup>2-</sup> ions. As explained earlier, it comes from the annihilation of positrons in the free volume defects of the Ag<sub>2</sub>O matrix, then in the interfaces and finally in the cationic vacancies within the nanocrystallites formed.

In order to explain it further, we note that the gradually rising trend of the peak heights of the CDBS ratio curves is indicative of the enhancing annihilation of positrons with the 2p electrons of oxygen. This is possible with the increasing formation of the CdMoO<sub>4</sub> phase when the concentration (x) of Ag<sub>2</sub>O is increased. Earlier, in a similar study by Bar *et al.* on the xCuI-(1-x)(0.5CuO-0.5MoO<sub>3</sub>) glass-nanocomposite systems with different values of x, it had been demonstrated that the

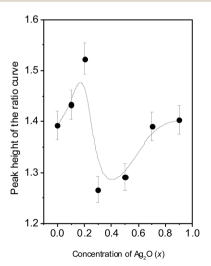


Fig. 13 The peak heights of the ratio curves *versus* concentration (x) of  $Ag_2O$  in the samples.

density of distribution of CuMoO<sub>4</sub> nanoparticles increased with the increase of CuI content.<sup>34</sup> A similar thermodynamic condition is prevalent in the present investigation too in which Ag<sub>2</sub>O is playing a strong catalytic role in the formation of the CdMoO<sub>4</sub> nanocrystallites and these in turn happen to act as very strong trapping centres for positrons. The discontinuity manifesting as an abrupt fall at x=0.2 is a result of translocation of the positron trapping sites from the Cd<sup>2+</sup>-monovacancies ( $V_{\rm Cd}^{6-}$ ) in the CdO nanocrystals<sup>35</sup> to the Mo<sup>6+</sup> monovacancies ( $V_{\rm Mo}^{6-}$ ) in the  $\alpha$ -MoO<sub>3</sub> (ref. 36 and 37) and then in the CdMoO<sub>4</sub> (ref. 38 and 39) nanocrystallites. While a reducing influence of the 2p electrons of O<sup>2-</sup> ions results into the fall in peak height at x=0.2, the enhanced formation of the CdMoO<sub>4</sub> nanocrystallites as evidenced from the increasing nanocrystallite size (Fig. 2) sustains the rising trend at further concentration (x) of Ag<sub>2</sub>O.

In this context, it may be useful to examine the structure of the different crystallites as the number and distance of the nearest neighbor O<sup>2-</sup> ions in each from a positron trapped at the metallic monovacancies may provide some more insight. Table 2 summarizes these parameters for the different crystalline phases present in the nanocomposites. CdO is essentially a face centred cubic structure and there are six neighboring  $O^{2-}$ ions octahedrally coordinated with each Cd2+ ion. The Cd-O bond length is 2.3-2.4 Å, as reported by several authors.8,35 An oxygen density parameter is calculated as the number of coordinated O2- ions divided by the volume of the regular octahedron formed by the Cd2+ and coordinated O2- ions. Similarly, Andersson et al.36 and Li et al.37 have described α-MoO3 as a distorted orthorhombic elementary cell with the Mo6+ ion octahedrally coordinated with six O2- ions. The structure of CdMoO<sub>4</sub> is an interesting one which crystallizes in a scheelite structure with each single crystal of CdMoO4 having eight symmetry elements and a body-centered orthorhombic primitive cell of two formula units.38,39 The Cd2+ ion is eightfold coordinated by O<sup>2-</sup> ions in the form of octahedron made up of compressed tetrahedrons of  $(MoO_4)^{2-}$ . The calculated oxygen density parameters are listed in Table 2. Although Ag<sub>2</sub>O has a simple cubic structure with each Ag1+ ion linearly coordinated with two O<sup>2-</sup> ions, one on either side, 40 we have included it also in the table by considering a fictitious octahedral volume around it.

The results indicate the probability for increasing annihilation events with the electrons of  $O^{2-}$  ions when the positrons are trapped in the  $Mo^{6+}$ -monovacancies instead of  $Cd^{2+}$ -monovacancies. The initial rise of the peak height in Fig. 13 is indicative of this. When  $CdMoO_4$  crystallites are formed, the initial trapping occurs in the  $Cd^{2+}$ -monovacancies of it. But, in this case, the oxygen density parameter is weakened and there is an abrupt fall of the peak height. With the positron trapping in the  $Mo^{6+}$ -monovacancies of the  $(MoO_4)^{2-}$ -tetrahedrons, the parameter increases again and the peak heights rise again. Positron trapping in  $Ag_2O$  oxide could have reversed the trend at the highest concentration of x=0.9 had the crystallites so formed been considerable in number. It should be noted further that the positron lifetime  $\tau_2$  in fact had shown an abrupt fall around x=0.1–0.2, then a small rising trend followed by

Table 2 The calculated oxygen density parameters for the different crystallite unit cells

Name of the nanocrystallite	Structure	Distance to the $O^{2-}$ -ion in octahedral coordination from the cation vacancy indicated in parenthesis $(a/2)$ (Å)	Volume of octahedron $v = a^3/6  (\mathring{A}^3)$	Number of coordinated $O^{2-}$ ions $(N)$	Oxygen density parameter $(N/\nu)$
CdO	Face centered cubic <sup>8,35</sup>	2.3-2.4 (Cd <sup>2+</sup> )	16.854	6	0.356
$\alpha$ -MoO <sub>3</sub>	Orthorhombic <sup>36,37</sup>	$1.85-2.35  (Mo^{6+}) \ge 2.10  (mean)$	12.348	6	0.486
$CdMoO_4$	Body centered	2.4-2.44 (Cd <sup>2+</sup> ) (octahedral)	18.897	8	0.423
	orthorhombic 11,38,39	1.75 (Mo <sup>6+</sup> ) (tetrahedral)	2.751	4	1.454
$Ag_2O$	Cubic <sup>40</sup>	2.04 (Ag <sup>1+</sup> ) (linear)	11.318 (fictitious octahedral)	2	0.177

a gradual decrease with increasing x (Fig. 9(a)), which clearly supported the above findings.

We also have calculated the S and W parameters that are popularly used in positron literature to comment on the qualitative nature of variation of the annihilation characteristics with the variational quantity chosen (x, in this case). These parameters are calculated according to their usual definitions that they respectively represent the fractions of low and high momentum electrons annihilated by the positrons. Accordingly, S is derived from the area-normalized counts within the central channels corresponding to 511  $\pm$  0.651 keV of the onedimensional projection of the CDB spectrum. For calculating the W parameter, equal number of channels on either side of  $p_{\rm L}$  $= 10 \times 10^{-3} m_0 c$  have been chosen, since this corresponded to the channel where the peak due to 2p electrons of O<sup>2-</sup> ions had appeared in the CDBS ratio curves (Fig. 12). The nature of variation of these two lineshape parameters also indicated the abrupt discontinuity at x = 0.2, as had been expected (Fig. 14). Also, by definition, the value of the S parameter is also a measure of the total defect volume within the samples. In this context, the striking similarity of variations of S with that of the free volume fraction  $f_V$  in Fig. 10 is notable although both positron lifetime and CDBS measurements are independent of each other and measured parameters are not directly correlated with each other.

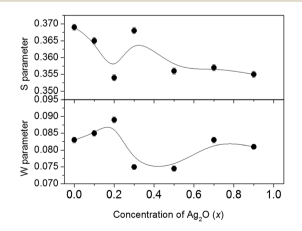


Fig. 14 The S and W parameters versus concentration (x) of  $Ag_2O$  in the samples.

# 4. Summary and conclusions

Metal oxide nanocomposites of  $xAg_2O-(1-x)(0.3CdO-0.7MoO_3)$ with x = 0.0 to 0.9 are investigated by, among other techniques, positron annihilation spectroscopy. The results of PAS essentially pointed towards the vacancy-type defects prevalent in the nanocomposite structure and their evolution under addition of Ag<sub>2</sub>O. The X-ray diffraction patterns were sharp indicating the presence of large crystallites but the TEM images confirmed the presence of nanocrystallites and thereby the nanocomposite nature. The optical band gaps obtained from the absorption spectra also showed systematic variation and indirectly indicated the growing influence of a new phase being formed. A distinct third positron lifetime component for samples with *x* > 0.3 clearly indicated the formation and annihilation of orthopositronium atoms within the free volume defects in the Ag<sub>2</sub>O amorphous matrix. Positrons were also trapped by the vacancytype defects within the nanocrystallites, the defects being Cd<sup>2+</sup>monovacancies in the initial stages of Ag2O incorporation and then the preferred trapping centres got successively shifted to metallic monovacancies of the other two nanocrystallites, viz., α-MoO<sub>3</sub> and CdMoO<sub>4</sub> at higher Ag<sub>2</sub>O concentrations. The latter inference is derived from the CDBS ratio curves in which variations of the characteristic peak heights indicated this trend. Besides, positrons also got annihilated in crystalline vacancies in the Ag<sub>2</sub>O matrix. There could be a rich concentration of oxygen vacancies, where positrons will not be trapped because of the positive charge of the vacancies.

# Acknowledgements

One of the authors (PMGN) wishes to thank Mr Aneesh Kumar S., Ms Drisya M. and Ms Sandhya C. of Payyanur College, Kannur, Kerala for their help in the positron annihilation experiments and data analysis during their summer project programme at Saha Institute of Nuclear Physics, Kolkata in January–February 2015. Mr Anish Karmahapatra and Ms Soma Roy of SINP, Kolkata are also gratefully acknowledged for their help in taking the XRD patterns and UV-Vis absorption spectra respectively. The authors are also grateful to Dr Biswarup Satpati of SINP, Kolkata for providing them the TEM images of the samples and some useful discussions. S. B. and R. K. wish to

Paper RSC Advances

acknowledge Council of Scientific and Industrial Research (CSIR), New Delhi for providing financial assistance, vide Sanction No. 03(1286)/13/EMR-II.

## References

- 1 B. Karmakar, *Glass Nanocomposites: Synthesis, Properties and Applications*, ed. Basudeb Karmakar, Klaus Rademann and Andrey Stepanov, Elsevier B.V., Amsterdam, 2016, ch. 1, pp. 3–53
- 2 A. R. Boccaccini, M. Erol, W. J. Stark, D. Mohn, Z. Hong and J. F. Manoe, *Compos. Sci. Technol.*, 2010, **70**, 1764–1776.
- 3 R. Krause-Rehberg and H. S. Leipner, *Positron Annihilation in Semiconductors Defect Studies*, Springer, Berlin, Germany, 1999, pp. 1–379.
- 4 P. M. G. Nambissan, *Nanotechnology: Synthesis and Characterization*, ed. Shishir Sinha, N. K. Navani and J. N. Govil, Studium Press LLC, Houston, U.S.A, 2013, ch. 16, vol. 2, pp. 455–491.
- 5 E. E. Assem, K. R. Mahmoud, T. Sharshar and C. Siligardi, *J. Phys. D: Appl. Phys.*, 2006, **39**, 734–739.
- 6 J. V. Olsen, P. Kirkegaard, N. J. Pedersen and M. Eldrup, *Phys. Status Solidi C*, 2007, **4**, 4004–4006.
- 7 A. Chatterjee, K. Ramachandran, S. Singh, S. S. Pande and M. D. Ghodgaonkar, *Proceedings of the DAE Symposium on Nuclear Physics*, 2002, 45, 145–146.
- 8 K. Usharani, A. R. Balun, V. S. Nagarethinam and M. Suganya, *Prog. Nat. Sci.: Mater. Int.*, 2015, 25, 251–257.
- 9 A. Stoyanova, R. Iordanova, M. Mancheva and Y. Dimitriev, *J. Optoelectron. Adv. Mater.*, 2009, **11**, 1127–1131.
- 10 M. M. Y. A. Alsaif, S. Balendhran, M. R. Field, K. Latham, W. Wlodarski, J. Zhen Ou and K. Kalantar-zadeha, Sens. Actuators, B, 2014, 192, 196–204.
- 11 Xi-E. Wang, T.-Y. Liu, Qi-R. Zhang, H.-Y. Zhang, M. Song, X.-F. Guo and Ji-G. Yin, Chin. Phys. Lett., 2008, 25, 1042–1044.
- 12 M. Guzik, E. Tomaszewicz, Y. Guyot, J. Legendziewicza and G. Boulonc, *J. Mater. Chem. C*, 2015, **3**, 4057–4069.
- 13 A. Patterson, Phys. Rev., 1939, 56, 978-982.
- 14 P. Manuel Diehm, P. Ágoston and K. Albe, *ChemPhysChem*, 2012, **13**, 2443–2454.
- 15 G. K. Williamson and W. H. Hall,  $Acta\ Metall.$ , 1953, 1, 22–31.
- 16 J. Tauc, Mater. Res. Bull., 1968, 3, 37-46.
- 17 M. Ghosh and C. N. R. Rao, *Chem. Phys. Lett.*, 2004, 393, 493-497.
- 18 R. M. Nieminen and M. J. Manninen, *Positrons in Solids Springer Series on Topics in Current Physics*, ed. P. Hautojarvi, Springer, Berlin, Germany 1979, vol. 12, ch. 4, pp. 145–195.

- 19 Y. C. Jean, K. Venkateswaran, E. Parsai and K. L. Cheng, *Appl. Phys. A*, 1984, 35, 169–176.
- 20 S. J. Tao, J. Chem. Phys., 1972, 56, 5499-5510.
- 21 M. Eldrup, D. Lightbody and J. N. Sherwood, *Chem. Phys.*, 1981, **63**, 51–58.
- 22 A. Uedono, R. Sadamoto, T. Kawano, S. Tanigawa and T. Uryu, J. Polym. Sci., Part B: Polym. Phys., 1995, 33, 891–897.
- 23 A. Seeger, J. Phys. F: Met. Phys., 1973, 3, 248-294.
- 24 I. K. MacKenzie, Positron Solid State Physics: International Summer School Proceedings Proceedings of the International School of Physics "Enrico Fermi", ed. W. Brandt and A. Dupasquier, North Holland, Amsterdam, 1983, pp. 196–264.
- 25 D. Wegner, J. Phys. F: Met. Phys., 1988, 18, 2291-2302.
- 26 J. M. Campillo Robles, E. Ogando and F. Plazaola, *J. Phys.: Condens. Matter*, 2007, **19**, 176222.
- 27 M. Fernández-Garcia and J. A. Rodriguez, *Nanomaterials: Inorganic and Bioinorganic Perspectives*, ed. C. M. Lukehart and R. A. Scott, Wiley, New York, 2008, pp. 453–512.
- 28 S. K. Sharma, P. K. Pujari, K. Sudarshana, D. Dutta, M. Mahapatra, S. V. Godbole, O. D. Jayakumar and A. K. Tyagi, *Solid State Commun.*, 2009, 149, 550–554.
- 29 A. Das, A. Chandra Mandal, S. Roy and P. M. G. Nambissan, J. Exp. Nanosci., 2015, 10, 622–639.
- 30 R. N. West, Adv. Phys., 1973, 22, 263-383.
- 31 R. W. Siegel, Annu. Rev. Mater. Sci., 1980, 10, 393-425.
- 32 P. Asoka-Kumar, M. Alatalo, V. J. Ghosh, A. C. Kruseman, B. Nielsen and K. G. Lynn, *Phys. Rev. Lett.*, 1996, 77, 2097– 2100.
- 33 Y. Nagai, T. Nonaka, M. Hasegawa, Y. Kobayashi, C. L. Wang, W. Zheng and C. Zhang, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **60**, 11863–11866.
- 34 A. K. Bar, D. Roy, R. Kundu, M. P. F. Graca, M. A. Valente and S. Bhattacharya, *Mater. Sci. Eng. B*, 2014, **189**, 21–26.
- 35 A. Bera, R. Thapa, K. K. Chattopadhyay and B. Saha, *J. Alloys Compd.*, 2015, **632**, 343–347.
- 36 G. Andersson and A. Magneli, *Acta Chem. Scand.*, 1950, 4, 793–797.
- 37 S. Li, W. B. Kim and J. S. Lee, *Chem. Mater.*, 1998, **10**, 1853–1862.
- 38 Y. Abraham, N. A. W. Holzwarth and R. T. Williams, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2000, **62**, 1733–1741.
- 39 M. Daturi, G. Busca, M. Madeleine Borel, A. Leclaire and P. Piaggio, *J. Phys. Chem. B*, 1997, **101**, 4358–4369.
- 40 A. Deb and A. K. Chatterjee, *J. Phys.: Condens. Matter*, 1998, **10**, 11719–11729.