Magnetization and optical bandgap of Cu-Mn vanadate-oxide mixed phase nanostructures

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Abstract Copper vanadate (CV) and manganese vanadate (MV) exhibit magnetic and optical properties that have drawn the attention. Due to CV polymorphism and phase multiplicity, CV is common to exist as mixed phases. In this study, nanostructures of mixed-phase CVs mixed with MV were synthesized hydrothermally followed by calcination at 400 °C, with Mn mole fractions 0.0, 0.4, 0.6, 1.0. The uncalcined and calcined Mn-Cu vanadates (MCVs) were investigated by XRD, SEM, TEM, FT-IR, EDX, ICP-AES, TGA, DTA, DSC, BET, XPS, and VSM. XRD analysis shows co-existence of multi-phase CVs with MnV₂O₆ and V oxides. Electron micrographs show nanostructures of multiple morphologies (rods, cubes, sheets, and irregular). As Mn content increased in the MCVs, their thermal stability increased, optical bandgap ($E_g$) declined from 2.46 to 1.60 eV, and magnetism diverted from the superparamagnetic-like to paramagnetic ($H_c$ from 1362 to 69 G and $M_r/M_s$ from 0.430 to 0.003). Magnetism parameters of calcined MCVs were more labile to Mn content variation compared to the uncalcined MCV counterparts.

Keywords Vanadate · Mixed oxides · Thermal stability · Band gap · Magnetism · Magnetisation · Superparamagnetic · Paramagnetic · Nanocomposites

Abbreviations
BET Brunauer–Emmett–Teller
$\Delta$ Difference in binding energy
cCV Calcined Cu vanadate
cMCV4 Calcined 0.4-Mn/0.6-Cu vanadate
cMCV6 Calcined 0.6-Mn/0.4-Cu vanadate
cMV Calcined Mn vanadate
CV Cu vanadate
DRS Diffuse reflectance spectroscopy
DSC Differential scanning calorimetry
DTA Differential thermal analysis
EDX Electron dispersive X-ray spectroscopy
$E_g$ Bandgap energy
eV Electronvolt
FT-IR Fourier transform infrared spectrum
Oe Oersted
Mₚ Saturation magnetization
MCV4 0.4-Mn/0.6-Cu vanadate
JCPDS Joint Committee on Powder Diffraction Standards
$H_c$ Coercivity
$\nu$ Photon energy

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ICP-AES  Inductively coupled plasma-atomic emission spectroscopy
MCV6  0.6-Mn/0.4-Cu vanadate
MCV  Manganese-copper vanadate
Mᵣ  Retentivity
MV  Mn vanadate
RT  Room temperature
SEM  Scanning electron microscopy
TEM  Transmission electron microscopy
TGA  Thermal gravimetric analysis
VSM  Vibrating sample magnetometer
xᵣ  Mn precursor mole fraction concerning Cu + Mn
XPS  X-ray photoelectron spectroscopy
XRD  X-ray diffraction
α  Absorption coefficient

Introduction

Transition metal divanadates (MV₂O₆ or M(VO₃)₂, M is a transition metal) have received great interest due to their application in various fields. Specifically, CuV₂O₆ nanoparticles have been applied in supercapacitors [1], batteries [2], photocatalytic water remediation [3], photoelectrocatalytic sensing [4], and photoelectrocatalytic catalysis [5]. Specifically, copper vanadate (CV) exhibits polymorphism and phase multiplicity, and takes the formulas CuV₂O₆, α-Cu₂V₂O₇ (Blossite), β-Cu₂V₂O₇ (Ziesite), Cu₃V₂O₈ (Pseudolynsite), γ-Cu₃V₂O₈ (McBirneyite), Cu₅V₂O₁₀, Cu₁₁V₆O₂₆ (Fingerite), and Cu₃(OH)₂V₂O₇(H₂O)₂ (Volborthite) [6]. Interestingly, the different CV phases can be viewed as mixtures of CuO with V₂O₅ in different ratios and mole fractions, as shown in Fig. 1 [7]. CVs can be synthesized with various approaches including co-precipitation [8, 9], hydrothermal [10, 11], thermal decomposition [6, 12], and solution combustion [7, 13].

Slight perturbation in the synthesis conditions leads to switching from one CV phase to another. Sometimes obtaining one pure phase can be a challenging task and just mixed-phase CVs originate [13–17]. The resulting mixed phases not only contain copper vanadates, but in some cases vanadium oxide may emerge as a distinct phase amongst the rest of the mixture. Hossain et al. and Keerthana et al. reported co-existence of V₂O₅ alongside with α-Cu₂V₂O₇ as a result of solution-combustion synthesis [3, 7].

Mixed-phase CVs have been synthesized, studied, and applied in various applications, such as photocatalytic water treatment [3], photoelectrocatalytic water treatment [13, 16], photoelectrocatalytic O₂ generation [18], photoelectrocatalytic water splitting [15]. Despite the controversy that may arise from the polymorphism and phase multiplicity of CVs, they have been pivotal in the aforementioned photoactivity-based studies, due to their visible-light absorptivity (E₉ 1.80–2.74 eV) [19], and magnetism investigations [20–26].

The present work aims to study the magnetization and optical bandgap of mixed-phase nanostructures of CVs with monoclinic Mn vanadate. As well as CVs, MnV₂O₆ exhibits visible-light photocactivity [27, 28] and magnetic properties [29–33]. The proposed Mn-Cu vanadate mixed-phase (MCV) was synthesized hydrothermally followed by calcination, with Mn mole fractions \( x_{\text{Mn}} = \frac{\text{Mn}}{\text{Cu} + \text{Mn}} \) of 0.0, 0.4, 0.6, 1.0. The investigated characteristics of the MCVs included their crystal structure, morphology, thermal stability, surface area, optical bandgap, and magnetization.

Experimental

Materials

Ammonium monovanadate extra pure (NH₄VO₃) (Oxford, India), manganese chloride tetrahydrate extra pure (MnCl₂.4H₂O) (Oxford, India), copper chloride

![Table 1](image-url)
dihydrate (CuCl₂·2H₂O) (Adwic Pharmaceutical and Chemicals Company, Egypt), and ethanol (C₂H₅OH) 99% (Adwic Pharmaceutical and Chemicals Company, Egypt). All chemicals and solvents were of analytical grade and used without further purification.

Hydrothermal synthesis of MCVs

MCVs of this study were synthesized according to a published method with some modifications [34]. Then, 4.8 mmol copper chloride (0.818 g) was dissolved into 80 mL of deionized water. Further, 9.6 mmol ammonium monovanadate (1.123 g) was dissolved into 80 mL deionized water at 60 °C, and then filtrated. Clear ammonium monovanadate solution was added slowly to the copper chloride solution under continuous vigorous stirring for 10 min. After the addition was completed, the resulting suspension was transferred into a 300 mL Teflon-lined stainless autoclave and maintained at 210 °C for 12 h. The autoclave was allowed to cool down to room temperature. The solid powder was collected by centrifugation, washed several times with deionized water and ethanol, and then dried at 60 °C for 4 h. The sample was labeled CV, then calcined at 400 °C for 2 h in the air (calcined sample is denoted as cCV).

Manganese vanadate was prepared by adding manganese chloride instead of copper chloride. The obtained manganese vanadate is labeled MV (before calcination) and cMV (after calcination). Mixed phases MCV4, MCV6, and their corresponding calcined samples cMCV4, cMCV6 were prepared by introducing different copper and manganese chlorides ratios as shown in Table 1.

Characterization methods and techniques

Powder x-ray diffraction (XRD) patterns were collected with BRUKER D 8 Advance, Germany. It operated with Cu Kα radiation wavelength 1.54°Å, at 40 kV, 40 mA, in the diffraction angle of 2θ from 5° to 80° at a scan rate of 5° min⁻¹. The Fourier transform infrared (FT-IR) spectra of samples before and after calcination was collected by Perkin Elmer spectrum version 10.5.3 IR spectrometer in the range 4000–400 cm⁻¹ at room temperature. The stoichiometric composition was obtained with inductively coupled plasma-atomic emission spectroscopy (ICP-AES) ICP-OES 5100 VDV, Agilent where 0.01 g of the MCVs after calcination were dissolved in 5 mL concentrated nitric acid then heated and completed with water to total 25 mL. Energy-dispersive X-ray spectroscopy (EDX) was conducted with field emission scanning electron microscope JEOL JSM 7000F with a 15 kV accelerating voltage. A 10 mm working distance was used to confirm the constituent elements to determine the elemental ratios. The size and morphology of the MCVs before calcination were characterized with (JEOL JEM-200CX2100F, Japan) transmission electron microscopy (TEM) with an accelerating voltage of 200 kV. Divanadate TEM samples were prepared by sonicating 0.01 g for 30 min with 5 mL of ethanol in a centrifugal tube. Cu grid 400 mesh with a thin carbon film (Quantifoil) was immersed into the dilute dispersion of samples. The morphology and microstructure of MCVs after calcination were analyzed using scanning electron microscopy (SEM) JSM-IT200. Thermal gravimetric analysis (TGA), differential thermal analysis (DTA), and differential scanning calorimetry (DSC) were carried out with SDT Q600 V20.9 Build 20; the MCV samples before calcination were put into the crucible and heated in air at a heating rate of 10 °C min⁻¹ from room temperature to final temperature 800 °C. UV–vis diffuse reflectance spectroscopy (DRS) was carried out at room temperature using a PerkinElmer Lamda-900 spectrophotometer in the range of

| Table 1 | Amounts of Cu and Mn precursors added during the synthesis of the Cu-Mn vanadate-oxide mixed phase nanostructures. xMn is Mn mole fraction with respect to Cu+Mn | | | | |
| Sample | xMn | CuCl₂·2H₂O | MnCl₂·4H₂O | pH of mixed reactants |
| | | mMol | Wt (g) | mMol | Wt (g) | |
| CV | 0.0 | 4.80 | 0.818 | 0.00 | 0.00 | 4.5 |
| MCV4 | 0.4 | 2.88 | 0.491 | 1.92 | 0.38 | 5.2 |
| MCV6 | 0.6 | 1.92 | 0.327 | 2.88 | 0.57 | 5.6 |
| MV | 1.0 | 0.00 | 0.00 | 4.80 | 0.95 | 6.1 |
200–800 nm. Brunauer–Emmett–Teller (BET) determined the specific surface areas of MCV samples after calcination by nitrogen adsorption at 77 K using a Belsorp-mini II (Japan). Before surface area analysis, the powders were degassed at 150 °C for 3 h under vacuum. To study the surface chemistry of MCV samples after calcination, X-ray photoelectron spectroscopy (XPS) was collected with K-ALPHA (Thermo Fisher Scientific, USA) of monochromatic X-ray Al K-alpha radiation (−10 to 1350 eV spot size 400 µm, at pressure 10−9 mbar, spectrum pass energy 200 eV, and at narrow-spectrum 50 eV). The magnetic properties of MCVs before and after calcination were measured with a vibrating sample magnetometer (VSM, Lake Shore-7410, USA) at room temperature with a magnetic field up to 20 kOe and magnetic moment sensitivity up to 1 m emu.

**Results and discussion**

**Characterization**

**X-ray diffraction**

Figure 2a and Table 2 of the virgin CV show peaks characteristic for three different phases. First, monoclinic volborthite Cu$_3$(OH)$_2$V$_2$O$_7$·2H$_2$O according to JCPDS Card No. 46–1443. Volborthite is a natural divanadate of formula Cu$_3$(OH)$_2$V$_2$O$_7$·2H$_2$O that exhibits interesting crystalline structure with porous framework [35]. As a metastable phase, Volborthite can be used as a precursor of another interesting mixed and layered crystalline phase composed of an octahedral copper-oxygen layer in coordination with tetrahedral vanadium layer. Second, monoclinic copper vanadium oxide Cu$_{0.261}$V$_2$O$_5$ according to pattern for Cu-Mn vanadate-oxide mixed phase nanostructures after calcination: (asterisk) Cu$_{564}$(V$_{19}$O$_{41}$)(OH)$_{9}$(H$_2$O)$_{11}$, (multiplication sign) V$_6$O$_{13}$, (plus sign) (NH$_4$)$_8$V$_{19}$O$_{41}$, (black circle) MnV$_2$O$_6$, (white circle) MnV$_2$O$_6$, (down-pointing triangle) V$_2$O$_5$.
The calcined samples in the air at 400 °C for 2 h, cCV, cMCV4, cMCV6, and cMV, show significant XRD changes than those of the virgin samples in addition to formation of new phases as shown in Fig. 2b and listed in Table 2. The XRD peaks of cCV in addition to formation of new phases as shown in cant XRD changes than those of the virgin samples 2 h, cCV, cMCV4, cMCV6, and cMV, show significant changes than those of the virgin samples at 2θ: 25.56°, which may be due to the presence of traces of orthorhombic V6O13 according to JCPDS Card No. 778–0983.

The monoclinic MnV2O6 (JCPDS Card No. 35–0139) was formed beside all previous phases; Cu3(OH)2V2O7·2H2O, Cu0.261V2O5, and V3O7. The highest noticed ratio is Cu0.261V2O5 as shown from the highest sharp peak at 2θ: 9.08°. In the MCV6, the peaks characteristic for Cu3.64(V4O11), V3O7 is noticed. Ammonium vanadium oxide hydroxide hydrate, (NH4)8(V19O41(OH)9)(H2O)11, exists as a trace amount with a characteristic peak at 2θ: 5.4° according to JCPDS Card No. 01–078-2016. In MV, the dominant phase is MnV2O6 beside a foreign peak at 2θ: 25.56°, which may be due to the presence of traces of orthorhombic V6O13 according to JCPDS Card No. 778–0983.

The XRD of cMCV6 demonstrates the presence of monoclinic MnV2O6 and V3O7 according to JCPDS Card No. 01–079-0226, Shcherbinaite V2O5, which is the only characteristic peak at 2θ: 5.4° of the uncalcined MCVs. It can be noticed that plenty of the bands of the uncalcined MCVs have disappeared upon calcination due to the elimination of H2O, CO2, and OH groups. The bands at 400–1010 cm−1 correlate to metal–oxygen bonds that have become stronger by calcination. Those at 795–1024 cm−1 are attributed to the vibrational mode of V=O bonds. The strong bands at 1016–1024 cm−1 of the V=O bond in V2O5 as a common product in all calcined samples. This band is very small for cMV due to the diminish of V2O5 [42]. The presence of the Mn–O bond could be observed at ~558 cm−1, which is closed to that reported in the literature at 564 cm−1 [39, 40]. The bands at 1401–1405 cm−1 of CO2 symmetric stretching-disappeared completely after calcination. The intensities of vibration bands at 1613–1620 cm−1 of the ν-OH stretching vibration and δH2O bending vibration of water molecules. The bands at 1016–1024 cm−1 are assigned to the ν-OH stretching vibration of water molecules [36].

Figure 3b shows the FTIR spectra of the calcined MCVs. It can be noticed that plenty of the bands of the uncalcined MCVs have disappeared upon calcination due to the elimination of H2O, CO2, and OH groups. The bands at 400–1010 cm−1 correlate to metal–oxygen bonds that have become stronger by calcination. Those at 795–1024 cm−1 are attributed to the vibrational mode of V=O bonds. The strong bands at 1016–1024 cm−1 of the V=O bond in V2O5 as a common product in all calcined samples. This band is very small for cMV due to the diminish of V2O5 [42]. The presence of the Mn–O bond could be observed at ~558 cm−1, which is closed to that reported in the literature at 564 cm−1 [39, 40]. The bands at 1401–1405 cm−1 of CO2 symmetric stretching-disappeared completely after calcination. The intensities of vibration bands at 1613–1620 cm−1 of the ν-OH stretching vibration and δH2O bending vibration of water molecules decreased due to water removal. The bands at 3000–2700 cm−1 are assigned to organic impurities decreased also. The bands at 3400–3600 cm−1 are assigned to the ν-OH stretching vibration of water molecules [36].

The EDX (Fig. 4) and ICP analysis show that the metals ratios (Cu, Mn, and V) are in good match with the starting concentrations of Cu2+, Mn2+, and VO3− added during the synthesis as shown in Table 3. Detailed raw EDX scan results are given in

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**Fourier transform infrared**

FT-IR spectra analysis was performed to investigate the structure and functional groups. Figure 3a shows the FTIR spectra of the uncalcined MCV samples. The bands observed at 500–1010 cm−1 correspond to the metal–oxygen bonds stretching and wagging (Cu–O, Mn–O, and V–O). The band at 899–886 cm−1 arises from the symmetric stretching vibration of ν3VO3, and that at 751 cm−1 is corresponding to the anti-symmetric stretching vibration of ν3VO3 of CV. The two bands at 530 cm−1 and 509 cm−1 are assigned to the ν3 symmetric and ν1 symmetric stretching modes of (V=O–V) of CV [36]. Those at 993–1010 cm−1 are corresponding to ν(V4+ = O) [37, 38]. The band observed at 575 cm−1 for MV is assigned for Mn–O vibration [39]. The bands at 1401–1405 cm−1 are assigned to symmetric stretching of CO2 due to the absorption of CO2 from the air [40]. The bands at 1613–1620 cm−1 correspond to the ν-OH stretching vibration and δH2O bending vibration of water molecules. The bands at 2700–3000 cm−1 are assigned to organic impurities on the surface of the samples (maybe resulting from handling) [41]. The bands at 3400–3600 cm−1 are assigned to the ν-OH stretching vibration of water molecules [36].

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**ICP and EDX elemental analysis**

The EDX (Fig. 4) and ICP analysis show that the metals ratios (Cu, Mn, and V) are in good match with the starting concentrations of Cu2+, Mn2+, and VO3− added during the synthesis as shown in Table 3. Detailed raw EDX scan results are given in
Table 2: Based on the XRD results, phases constituting the uncalcined and calcined Cu-Mn vanadate-oxide mixed phase nanostructures, in addition to their average particle sizes estimated with the Scherrer equation. P: present, M: major and T: traces. JCPDS Card numbers: (a) 46–1443, (b) 01–079-0796, (c) 20–1378, (d) 35–0139, (e) 778–983, (f) 01–078-2016, (g) 1–085–0226, (h) 9–0387 and (i) 45–1054.

| Phase                                      | CV   | MCV4 | MCV6 | MV   |
|--------------------------------------------|------|------|------|------|
| Cu$_3$(OH)$_2$V$_2$O$_7$·2H$_2$O            | P    | P    | –    | –    |
| Cu$_{0.261}$V$_2$O$_5$                      | P    | P    | P    | –    |
| V$_3$O$_7$                                 | T    | P    | P    | –    |
| MnV$_2$O$_6$                               | –    | P    | P    | M    |
| V$_3$O$_{12}$                              | –    | –    | –    | T    |
| (NH$_4$)$_8$(V$_{19}$O$_{41}$(OH)$_9$)(H$_2$O)$_{11}$ | –    | –    | T    | –    |
| Average $D_{Sch}$ (nm)                      | 25   | 15   | 19   | 20   |

| Phase                                      | cCV  | cMCV4 | cMCV6 | cMV  |
|--------------------------------------------|------|-------|-------|------|
| Cu$_{1.64}$(V$_4$O$_{11}$)                 | P    | P     | P     | –    |
| V$_3$O$_5$                                 | P    | P     | P     | T    |
| CuV$_2$O$_6$                               | P    | P     | –     | –    |
| MnV$_2$O$_6$                               | –    | P     | P     | T    |
| Average $D_{Sch}$ (nm)                      | 13   | 20    | 19    | 27   |

Figure 5 is the TEM micrographs of the synthesized MCVs before calcination. CV shows nanowire bundles with several micrometers in length and cubic nanostructures of an average size of 100–300 nm. MCV4 shows nanowires with a slight increase in diameter (around 50 nm), cubic structures, and nanosheets (around 500 nm size). Since MV TEM images show all nanosheets only, these nanosheets are assumed to be of the MnV$_2$O$_6$ phase. MCV6 shows an increase in the abundance of MnV$_2$O$_6$ nanosheets and the thickness of nanowires, while absence of the cubic structures. MV shows MnV$_2$O$_6$ nanosheets of different sizes.

Fig. S1, S2, S3, and S4 in the supplementary information for samples cCV, cMCV4, cMCV6, and cMV, respectively.

Morphology of the uncalcined and calcined copper/manganese vanadates by scanning and transmission electron microscopy

Fig. 3 FTIR spectra of Cu-Mn vanadate-oxide mixed phase nanostructures before a and after b calcination.
Figure 6 shows SEM micrographs of cMCVs. cCV consists of nano-rods beside irregular shapes. This may be attributed to the presence of multi-phases of copper vanadates and \( \text{V}_2\text{O}_5 \). In cMCV4, shapes are more irregular, while in cMCV6, sheets the abundance with fewer rods than cCV. In cMV, there are no rods, and the sample is all nanosheets either stacked or separate.

Table 3 The Cu, Mn, and V elemental ratios according to the ICP and XPS techniques, in addition to the specific surface area, average pore volume, and pore diameter according to the BET technique, all for the calcined Cu-Mn vanadate-oxide mixed phase nanostructures. (*) The ideal stoichiometric ratio of \( \text{V}/(\text{Cu}+\text{Mn}) \) based on the expected formula \( \text{MV}_2\text{O}_6 \) should be 2, as shown in the brackets. (#) According to the added ratios during the synthesis, \( x \) was expected to change as 0.00, 0.40, 0.60, and 1.00 for samples cCV, cMCV4, cMCV6, and cMV, respectively, as shown in the brackets.

|          | cCV   | cMCV4 | cMCV6 | cMV   |
|----------|-------|-------|-------|-------|
| **ICP**  |       |       |       |       |
| Cu       | 0.27  | 0.13  | 0.06  | 0.00  |
| Mn       | 0.00  | 0.08  | 0.19  | 0.34  |
| V        | 0.73  | 0.79  | 0.75  | 0.66  |
| \( \text{V}/(\text{Cu}+\text{Mn}) \)* | 2.70 (2.00) | 3.76 (2.00) | 3.00 (2.00) | 1.94 (2.00) |
| \( x = \text{Mn}/(\text{Mn}+\text{Cu}) \)# | 0.00 (0.00) | 0.38 (0.40) | 0.76 (0.60) | 1.00 (1.00) |
| **XPS**  |       |       |       |       |
| Cu       | 0.25  | 0.06  | 0.04  | 0.00  |
| Mn       | 0.00  | 0.04  | 0.04  | 0.16  |
| V        | 0.75  | 0.90  | 0.92  | 0.84  |
| \( \text{V}/(\text{Cu}+\text{Mn}) \)* | 3.00 (2.00) | 9.00 (2.00) | 11.50 (2.00) | 5.25 (2.00) |
| \( x = \text{Mn}/(\text{Mn}+\text{Cu}) \)# | 0.00 (0.00) | 0.40 (0.40) | 0.50 (0.60) | 1.00 (1.00) |
| **BET**  |       |       |       |       |
| \( S_{\text{BET}} \) (m\(^2\)/g) | 4.7   | 8.6   | 5.4   | 1.0   |
| Pore volume (cm\(^3\)/g) \( \times 10^{-2} \) | 6.13  | 11.60 | 7.96  | 2.30  |
| Main Pore diameter (nm) | 52.3  | 53.9  | 59.4  | 92.7  |
Thermogravimetric analysis of copper and manganese vanadates (TGA, DTA, and DSC)

Figure 7 and Table 4 show the thermal analyses of the uncalcined MCVs carried out under an air atmosphere at a heating rate of 10 °C/min from the ambient temperature to 800 °C. The weight losses observed in the TG thermograms (Fig. 7a) are generally due to the removal of adsorbed water, crystallization water, and hydroxyl groups. The results are summarized in Table 4. For CV, the loss in weight takes place at four stages: the first stage starts from ambient temperature to 175 °C, the weight loss is 5.5% which may be due to the removal of adsorbed water and some crystallization water of the Volborthite phase. In the second stage, starting from 175 to 260 °C, the weight loss is 4.8% which is attributed to the complete removal of crystalline water of volborthite. In the third stage, starting from 260 to 400 °C, the weight loss is 3.9% due to removing two hydroxyl groups of Volborthite [43, 44]. The TG curve also shows another weight loss started at about 635 °C, which corresponds to the melting point of CuV2O6 in the CV sample. An endothermic peak accompanying this step appeared in each DTA and DSC thermogram (Fig. 7b and c). In the MCV4 sample, the weight loss of the melting step started at about 698 °C due to the melting point of V2O5 [45]. With decreasing the ratio of copper in the investigated samples, the weight loss also decreases since the samples contain fewer phases of crystallization water and hydroxyl groups (as pointed out in the XRD analysis and Table 2). MV exhibits minimal weight loss, which may be due to its phase stability and it shows only phase of MnV2O6 all over the thermal treatment.

Fig. 5 TEM images of the uncalcined Cu-Mn vanadate-oxide mixed phase nanostructures
Fig. 6  SEM images of the calcined Cu-Mn vanadate-oxide mixed phase nanostructures. The scale bar is equivalent to 1 µm

Fig. 7  Plots of TG (a), TDA (b), and DSC (c) of the uncalcined Cu-Mn vanadate-oxide mixed phase nanostructures
The optical spectra of the MCVs, before and after calcination, are shown in Fig. 8a and b. All MCVs show two absorption peaks; a small one at the UV range and broadband in the visible region. The results are summarized in Table 5. The bands that appeared in UV and visible regions are attributed to several phases of copper vanadates accompanying MnV₂O₆ and V₂O₅.

The optical bandgaps of the MCVs in the visible region were determined using Tauc equation as shown in Fig. 8c and d.

\[
(\alpha h\nu)^n = B(h\nu - E_g)
\]

where \( h\nu \) is the photon energy (\( E_{\text{photon}} \)), \( \alpha \) is the absorption coefficient, is a proportionality constant, and “\( n \)” depends on the type of transition, \( n = 1/2 \) for direct transmission, and \( n = 2 \) for indirect transmission [46]. The bandgap energy (\( E_g \)) was determined by extrapolating the linear part of \((\alpha h\nu)^2 \) vs. \( h\nu \) plot to intercept the horizontal \( E_{\text{photon}} \) axis as shown in Fig. 8c and d. The calculated \( E_g \) values for calcined and uncalcined MCVs are given in Table 5 and plotted versus \( x_{\text{Mn}} \) in Fig. 8e. In both uncalcined and calcined MCVs, \( E_g \) declined with the increase of \( x_{\text{Mn}} \), whereas the uncalcined ones were more responsive.

### Specific surface area by BET

The N₂ adsorption–desorption isotherms, as shown in Fig. 9, are identified as V type, which exhibits low adsorption at low gas pressure, indicating relatively weak affinity between gas and solid [47]. The sharply upward hysteresis loops of the samples close to \( P/P_0 = 1 \) suggest that the adsorption occurs in micropores and/or mesopores, where the interactions between the adsorbent and the adsorbate are relatively weak. The surface data of the specimens investigated are listed in Table 3. The surface area increases in the sequence cMCV4 > cMCV6 > cCV > cMV. The minimal surface area of cMV is attributed to the nanosheet structure, which is subject to stacking. One the other hand, cMCV4 and cMCV6 act show maximal surface area, probably due to the multiplicity of their phase composition. The pore size exhibits a broad range within 52–93 nm. These results further confirm the presence of macropores [48].

### X-ray photoelectron spectroscopy

The XPS spectra of cCV, cMCV4, cMCV6, and cMV, and their analysis are given in Fig. 10 and Table 6, respectively. The figure shows the presence of Cu, Mn, V, and O, and absence of any other foreign elements referring to the purity of the samples, which is further confirmed via XRD, FTIR, ICP, and EDX techniques. Due to the spin–orbit coupling, there are peaks for \( 2p_{1/2} \) at lower energy than those of \( 2p_{3/2} \), for each of Cu, Mn, V. Each of the three elements exist in the samples in two oxidation states, one higher than the other, which result in two peaks under each of \( 2p_{1/2} \) and \( 2p_{3/2} \). However, the high and low valency peaks tend to overlap and hide under the parent peaks of \( 2p_{1/2} \) and \( 2p_{3/2} \), and in order to show them in high resolution, those peaks are deconvoluted by the XPS peak differentiation-imitation method. The peak at \( \sim 284.8 \) eV is characterized by C 1 s (called adventitious carbon) and originates from the adventitious contamination layer [49]. C 1 s peak

| Table 4 Thermogravimetric data of uncalcined Cu-Mn vanadate-oxide mixed phase nanostructures |
|-----------------|------------------|------------------|
| Temperature (°C) | Weight loss (%) | Significance |
| CV 27–175        | 5.5              | Removing adsorbed water, besides the elimination of crystallized water molecules |
| 175–260          | 4.8              | |
| 260–400          | 3.9              | |
| 635–800          | 1.3              | |
| MCV4 27–170      | 3.4              | Removing adsorbed water, besides the elimination of crystallized water molecules |
| 170–390          | 5.1              | |
| 390–600          | 0.7              | |
| 600–698          | 0.9              | |
| 698–760          | 1.7              | |
| MCV6 27–175      | 2.3              | Removing adsorbed molecules |
| 175–410          | 2.7              | |
| MV 27–550        | 2.5              | Removing adsorbed water |
Fig. 8 UV–vis absorption spectra of the Cu-Mn vanadate-oxide mixed phase nanostructures, before (a) and after (b) calcination. The Tauc plots [(αhν)^2 vs. hν] before (c) and after (d) calcination. Dependence of bandgap of the uncalcined MCVs (blue curve in inset e) and calcined MCVs (red curve in inset e) on x_{Mn}.

is used as a reference to obtain the exact peaks values [50, 51]. The intensity of the carbon peak increased as the amount of time spent inside the instrument increased [52].
XPS spectra of Cu\(^{2+}\) have characteristic satellite peaks, which are prominent at 940.7, 943.3, 961.7 eV; these satellite peaks are characteristic of materials having ground-state d\(^9\) configuration, i.e., Cu\(^{2+}\) [53, 54]. The binding energy difference between 2p\(^{3/2}\) and 2p\(^{1/2}\) is 19.95 eV, which is close to the literature slandered value [55, 56]. The molar ratio between Cu\(^{1+}\) and Cu\(^{2+}\) are nearly given Table 6, based on 2p\(^{1/2}\) peaks analysis. In the manner of Cu analysis, Mn shows 2 main peaks (2p\(^{3/2}\) and 2p\(^{1/2}\)) and each of them is deconvoluted to two sub-peaks for Mn\(^{2+}\) and Mn\(^{3+}\) [57, 58]. The table also shows the molar ratio between Mn\(^{2+}\) and Mn\(^{3+}\) according to the 2p\(^{1/2}\) peaks analysis. The peaks appearing around 516.3–516.78 eV and 521.99–524.32 eV are assigned to the V 2p\(^{3/2}\) and 2p\(^{1/2}\). Due to the multivalency of V (V\(^{3+}\), V\(^{4+}\), and V\(^{5+}\)), multiple sub-peaks appear in the deconvolution of the high resolution [59–62]. The second highest-binding-energy peak is found at 531.53 eV (perhaps a third peak at 533.00 eV). The O 1s XPS signals are divided into three peaks for cCV, cMCV\(_6\), cMCV\(_4\), and four peaks for cMCV4. The prominent peak in all samples has binding energy in the range of 529.09–529.93 eV, which is usually for lattice O of several spinel 3d metal oxides [63]. However, the exact assignment of the higher binding energy peaks is somewhat complex and controversial as numerous factors like surface defects, contaminants, impurities, or chemisorbed oxygen species could result in the appearance of shoulder peaks [64, 65].

**Magnetic properties**

Figure 11 shows the M-H magnetization of the uncalcined and calcined MCVs under the influence of an external magnetic field at room temperature, and Table 7 summarizes their magnetism parameters. Figure 12 represents the impact of \(x_{\text{Mn}}\) on the various magnetism parameters, including coercivity field (\(H_c\)), magnetization saturation (\(M_s\)), remanent magnetization (\(M_r\)), and remanence ratio (\(M_r/M_s\)).

According to Fig. 12, all four parameters are more responsive to the Mn/Cu ratio in the calcined MCVs than in their uncalcined counterparts. Furthermore, with the increase of Mn, both \(H_c\) and \(M_s\) seem to change linearly in the uncalcined samples, while the former declines and the later rises. \(M_r\) does not show a remarkable response to the variation in \(x_{\text{Mn}}\) in the uncalcined MCVs. In the calcined MCVs, all of \(H_c\), \(M_s\), and \(M_r/M_s\) decrease remarkably with the increase of \(x_{\text{Mn}}\), unlike \(M_r\), which increases. At \(x_{\text{Mn}} > 0.6\) in the calcined MCVs, none of \(H_c\), \(M_s\) or \(M_r/M_s\) is responsive to the changes in \(x_{\text{Mn}}\). As \(x_{\text{Mn}}\) increased, the MCVs tend to divert from the superparamagnetic-like to paramagnetic (especially the calcined), where the uncalcined \(H_c\) declined from 396 to 118 G and \(M_r/M_s\) from 0.110 to 0.006 and for the calcined from 1362 to 69 G and from 0.430 to 0.003, respectively.

![Fig. 9 N\(_2\) adsorption/desorption BET isotherms of the calcined Cu-Mn vanadate-oxide mixed phase nanostructures](image)

| Table 5 Absorption peaks wavelengths in the UV and visible spectra as well as \(E_g\) values of the uncalcined and calcined Cu-Mn vanadate-oxide mixed phase nanostructures |
|-----------------------------------|--------|--------|--------|--------|
|                                   | CV     | MCV4   | MCV6   | MV     |
| UV peaks (nm)                     | 215    | 240    | 234    | 238    |
| Visible peaks (nm)                | 380    | 382    | 380    | 516    |
| Band gap \((E_g)\) (eV)           | 2.46   | 1.98   | 1.94   | 1.60   |
|                                   | cCV    | cMCV4  | cMCV6  | cMV    |
| UV peaks (nm)                     | 232    | 240    | 235    | 237    |
| Visible peaks (nm)                | 380    | 348    | 380    | 545    |
| Band gap \((E_g)\) (eV)           | 2.37   | 2.13   | 2.10   | 1.63   |
The magnetic characteristics of the studied materials depend on the magnetic interaction (superexchange interaction) between metal ions with magnetic moments in the crystal lattice. Thus,

Fig. 10 XPS survey scans for the calcined Cu-Mn vanadate-oxide mixed phase nanostructures (top panel) and their corresponding Cu, Mn, V, and O high-resolution peaks (the bottom 10 panels)
VSM studies were performed at room temperature within −20 kOe to +20 kOe. As shown in Fig. 11 and 12, the uncalcined MCVs show weak hysteresis with a very small magnetic coercivity at a low magnetic field, indicating superparamagnetic-like behavior. The value of magnetic coercivity of the uncalcined MCVs follows the order: CV > MCV4 > MCV6 > MV, as shown in Table 7 and Fig. 12.

It is noted that all the studied MCVs did not reach complete saturation even under a high magnetic field of 20 kOe. Several authors have reported the reduction of magnetization in NPs and proposed mechanisms to explain the no-saturation behavior in a high magnetic field. It can be attributed to the presence of a spin disordered surface layer, which requires a larger magnetic field to reach saturation magnetization [66]. The saturation magnetization (estimated from the linear extrapolation of \(M_\text{vs. } 1/H\) plot) of all samples is listed in Table 7. The chemical composition can significantly influence the magnetization behavior because of changes in the distribution of cations and the particle size. The reduction of the magnitude of magnetization is ascribed to the noncollinear spin arrangement at the particle surface and the difference in the magnetization characteristic of two sub-lattices due to cation redistribution [67]. The disordered or misaligned surface spins weaken the total magnetization of the material NPs, with small retentivity and coercivity values. These values indicate that the thermal variations are enough to dominate the anisotropic energy barrier of the studied samples and reverse the magnetization direction spontaneously. As seen in Table 7 and Fig. 12, the retentivity (\(M_r\)) and coercivity (\(H_c\)) values show variation concerning sample composition, which can be attributed to the interaction among the oxygen and metal ions in the sublattice of the crystals.

**Conclusion**

Due to CV polymorphism and phase multiplicity, the Cu, Mn, and V stoichiometry in the MCVs did not follow the metal divanadate formula \(MV_2O_6\) (\(M = \text{Cu}\) or \(\text{Mn}\)) nor did it follow the precursor Mn mole fraction \(x_{\text{Mn}} = \frac{\text{Mn}}{\text{Cu}+\text{Mn}} = 0.0, 0.4, 0.6, 1.0\). The \(\frac{V}{\text{Cu}+\text{Mn}}\) ratio reached as high as 3.0–11.5, while it was expected to

| Table 6 | XPS-based core-electrons binding energies (eV) of Cu, Mn, V, and O constituents in the calcined Cu-Mn vanadate-oxide mixed phase nanostructures |
|---------|------------------------------------------------------------|
|         | cCV            | cMCV4         | cMCV6         | cMV           | References |
| Cu      |                |               |               |               |            |
| 2p^{3/2}| 931.68 (Cu^{1+}) | 931.18 (Cu^{1+}) | 931.33 (Cu^{1+}) | –             | [53–56]    |
|         | 934.41 (Cu^{2+}) | 933.86 (Cu^{2+}) | 933.78 (Cu^{2+}) |               |            |
| 2p^{1/2}| 951.52 (Cu^{1+}) | 950.85 (Cu^{1+}) | 951.01 (Cu^{1+}) | –             |            |
|         | 954.36 (Cu^{2+}) | 953.76 (Cu^{2+}) | 953.86 (Cu^{2+}) |               |            |
| Satellite peaks | 940.74, 943.26, 961.7 | 940.14, 942.92, 961.22 | 940.26, 943.35, 961.06 | –             |            |
| Cu^{1+}:Cu^{2+} | 0.16:0.84 | 0.24:0.76 | 0.21:0.79 |               |            |
| Mn      |                |               |               |               |            |
| 2p^{3/2}| –             | 640.73 (Mn^{2+}) | 640.08 (Mn^{2+}) | 640.83 (Mn^{3+}) | [57, 58] |
|         |               | 641.33 (Mn^{3+}) | 642.12 (Mn^{3+}) |               |            |
| 2p^{1/2}| –             | 652.41 (Mn^{2+}) | 652.71 (Mn^{2+}) | 652.44 (Mn^{3+}) |            |
|         |               | 656.27 (Mn^{3+}) | 653.30 (Mn^{3+}) |               |            |
| Satellite peaks | –         | 644.72, 648.42 | 645.04, 647.96 | 646.99       |            |
| Mn^{2+}:Mn^{3+} | 1.00:0.00 | 0.18:0.82 | 0.29:0.71 |               |            |
| V       |                |               |               |               |            |
| 2p^{3/2}| 516.3, 516.66, 516.74 | 516.43, 516.44, 516.62 | 516.33, 516.6, 516.78 | 516.77, 516.99, 517.64 | [59–62]    |
| 2p^{1/2}| 523.09, 524.32 | 523.00, 524.12 | 522.91, 524.22 | 521.99, 524.11, 525.19 |            |
| O       | 529.09, 529.79, 530.24 | 529.45, 530.69, 531.8, 532.59 | 529.57, 529.95, 532.03 | 529.93, 530.75, 531.9 |            |
be only 2 if it followed the divanadate formula MV₂O₆.

Regardless of the precursor Cu and Mn mole fractions and regardless of the post-synthesis thermal treatment, each MCV sample consisted of 2–4 different vanadate-oxide phases of (NH₄)₈(V₁₉O₄₁(OH)₉)(H₂O)₁₁, V₂O₅, V₃O₇, V₆O₁₃, Cu₃(OH)₂V₂O₇·2H₂O, Cu₀.₂₆₁V₂O₅, Cu₃.₆₄V(V₄O₁₁), CuV₂O₆, or MnV₂O₆. When there was no copper, vanadium oxide was a trace phase accompanying the major phase of MnV₂O₆. Phase multiplicity of the MCVs clearly appears in the electron micrographs, where particles morphology in each MCV varies between nanowires, bundles of nanowires, nanocubes, nanosheets, and irregular structures.

![Fig. 11](image-url)

**Fig. 11** M–H magnetization plots of the uncalcined Cu-Mn vanadate-oxide mixed phase nanostructures (insets a–h) and the calcined ones (insets i–p). Plots b, d, f, h, j, l, n, and p are zoomed-in M–H plots for clarification of the hysteresis behavior.
As Mn content increased, the uncalcined MCVs showed more thermal stability and their optical bandgap remarkably declined. The calcined MCVs showed less bandgap response to the Mn-content variation than the uncalcined ones, although they all lay within 1.60–2.46 eV (all visible).

The uncalcined MCVs tend to divert from the superparamagnetic-like to be paramagnetic with the increase of Mn content, since both $H_c$ and $M_s$ seem to linearly decline and rise, respectively. This behavior is more evident in the calcined MCVs, where all of $H_c$, $M_s$, and $M_r/M_s$ decrease remarkably with the increase of $x_{Mn}$, unlike $M_s$, which increases.

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**Data availability** The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Prime novelty statement** All data and work included in this paper are original and have not been published or

| Table 7 Magnetism parameters of the uncalcined and calcined Cu-Mn vanadate-oxide mixed phase nanostructures |
|-------------------------------------------------|----------------|----------------|----------------|
| CV  | MCV4 | MCV6 | MV |
| $H_c$ (G) | 396 | 272 | 249 | 118 |
| $M_s$ (emu/g) | 0.07 | 0.42 | 0.77 | 1.24 |
| $M_r$ (emu/g × 10^-3) | 8.0 | 6.9 | 13.0 | 7.6 |
| $M_r/M_s$ (× 10^-3) | 110 | 17 | 15 | 6 |
| $H_c$ (G) | 1362 | 1277 | 140 | 69 |
| $M_s$ (emu/g) | 0.51 | 0.65 | 0.80 | 1.23 |
| $M_r$ (emu/g × 10^-3) | 220 | 110 | 7 | 4 |
| $M_r/M_s$ (×10^-3) | 430 | 170 | 9 | 3 |

**Fig. 12** Plots of various magnetism parameters $H_c$ (a), $M_s$ (b), $M_r$ (c), and $M_r/M_s$ (d) versus Mn molar ratio in the synthesis precursors added during the synthesis of the Cu-Mn vanadate-oxide mixed phase nanostructures. The blue and red curves represent the uncalcined and calcined samples, respectively.
under consideration of publishing by any other periodical or publisher.

**Declarations**

**Competing interests** The authors declare no competing interests.

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