RSC Advances



PAPER



Cite this: RSC Adv., 2020, 10, 10491

Cesium polytungstates with blue-tint-tunable near-infrared absorption

Satoshi Yoshio, (10 ** Masao Wakabayashi ** and Kenji Adachi (10 **)

Revisiting Wöhler's method (1824), Cs-doped tungsten bronzes were synthesized by reducing Cspolytungstate at high temperature, and were pulverized into nanoparticles for determining their optical properties. The high-temperature reduced Cs₄W₁₁O₃₅ crystals absorbed strongly in the near-infrared, providing an improved luminous transparency with a less-bluish tint than normal $Cs_{0.32}WO_{3-\nu}$ synthesized in a reductive atmosphere. The high-temperature reduction caused an orthorhombic-tohexagonal phase transformation and a nonmetal-metal transition, which was monitored by spectrophotometry, X-ray diffraction, and X-ray photoelectron spectroscopy measurements, assisted by a first-principles analysis using a DFT+U method. The high-temperature reduction of $Cs_4W_{11}O_{35}$ is concluded to decrease the number of W deficiencies and produce oxygen vacancies, releasing both free and trapped electrons into the conduction band and thereby activating the near-infrared absorption. The comparatively narrow bandgap of $Cs_4W_{11}O_{35}$ was identified as the origin of the less-bluish tint of the produced Cs tungsten bronzes.

Received 17th January 2020 Accepted 3rd March 2020

DOI: 10.1039/d0ra00505c

rsc.li/rsc-advances

Introduction

Recently, hexagonal tungsten bronze (HTB) nanoparticles have demonstrated a high-level compatibility of luminous transparency and near-infrared (NIR) absorption, 1-3 prompting their application in automotive and architectural windows, laser welding of resins, cancer therapies, 4,5 and related technologies.

The origin of the strong NIR absorption in reduced tungsten oxides and tungsten bronze nanoparticles has been attributed to localized surface plasmon resonance (LSPR) of free electrons and polaronic excitation of trapped electrons. 6-16 In Cs-doped HTB (Cs-HTB), recent analyses indicate that W-5d orbitals in the lower conduction band are occupied by the free and trapped electrons originating from Cs⁺ and oxygen vacancies (V_O), respectively. 17,18,40 The Vos have been analyzed to play a major role in the LSPR and polaronic excitation. 19,20 On the other hand, the visible transparency of Cs-HTB nanoparticles, which is highest among the HTBs, has not been considered in detail, and its origin remains unclear. In addition, the slight bluish tint accompanying the transmitted light must be resolved in actual applications.

The imaginary part ε_2 of the dielectric function, which represents the absorption of a photon by an electron, is small at visible frequencies in Cs-HTB (see Fig. 1). The small ε_2 is caused mainly by a wide band gap. Although the band gap of Cs-HTB is

below 3.3 eV, the calculated electronic structure of Cs-HTB^{17,18} suggests that Fermi's selection rule prohibits certain electronic transitions such as W-5d \rightarrow W-5d and O-2p \rightarrow O-2p, which are involved in the hybridized W-5d/O-2p orbitals. This prohibition of certain transitions between metal-d and ligand-p orbitals further amplifies the visible transparency, enabling it even when the conduction and valence bands overlap in specific directions, as occurs in LaB₆.21 The optical absorption of Cs-HTB nanoparticles is known large at both frequency ends of the visible region. On the blue side of the spectrum, the ε_2 is enlarged by band-edge interband transition; on the red side, strong absorption occurs by LSPR and polaronic electronic

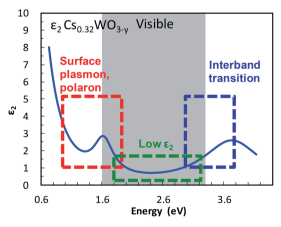


Fig. 1 Transparency and color-determining factors shown in the imaginary component ε_2 of the dielectric function of Cs_{0.32}WO_{3-y}.

^aDepartment of Computer-Aided Engineering, Sumitomo Metal Mining Co., Ltd., Ehime 792-0001, Japan. E-mail: satoshi.yoshio.w5@smm-g.com

^bIchikawa Research Center, Sumitomo Metal Mining Co., Ltd., Ichikawa, Chiba 272-8588, Japan

transition.^{13,20} Therefore, the blue-tint problem in Cs-HTB nanoparticles reduces to the problem of controlling the band gap and the conduction-band electrons.

Cs-HTB is usually prepared by the solid-phase reaction in a reductive atmosphere. This study adopts a different approach that controls the band gap and V_O-derived electrons. Magnéli and Blomberg²² summarized various methods of obtaining tungsten bronzes. The reduction of polytungstates (Cs₂O·nWO₃) in a steam of hydrogen gas flow was originally reported by Wöhler²³ in 1824, but has been scarcely followed thereafter, so the synthesis product has not been reported in detail. Starting from a charge-neutral polytungstate (the Cs₄W₁₁O₃₅ phase), we here synthesize a series of Cspolytungstates (referred to as CPT) with a significantly reduced blue tint by the Wöhler's method. This reduction process involves the orthogonal-to-hexagonal phase transformation as well as the nonmetal-metal transition. The change in optical absorption is then related to these phase transitions.

II Experimental

II-1 Materials

Raw materials Cs₂CO₃ and WO₃ of 99.9% purity were purchased from Fujifilm Wako Pure Chemical Corporation (Tokyo, Japan), and methyl isobutyl ketone (MIBK) of 99.5% purity was purchased from Kanto Chemical Co., Inc. (Tokyo, Japan).

As the starting material, $Cs_4W_{11}O_{35}$ was selected because its Cs/W ratio was close to that of $Cs_{0.32}WO_{3-y}$. $Cs_4W_{11}O_{35}$ was prepared by heating the Cs_2CO_3 and WO_3 raw materials in air at 800 °C for 30 hours or at 850 °C for 5 hours, respectively. The X-ray diffraction (XRD) powder pattern revealed the primary $Cs_4W_{11}O_{35}$ phase with a minutely mixed $Cs_6W_{11}O_{36}$ phase.

II-2 Powder and dispersion preparation

In the conventional gas flow reduction (GFR) method¹⁹ or the vacuum encapsulation method,²⁴ the crystallization of $Cs_{0.32}$ - WO_{3-y} and oxygen reduction proceed in parallel with the high-temperature reduction. On the other hand, when polytungstate crystals are reduced at high temperature, high crystallinity is expected from the early stages of reduction, which was considered as an advantage of the Wöhler's method. Thus, $Cs_4W_{11}O_{35}$ was heated to 700–900 °C in an Ar gas flow in a tubular furnace. While maintaining the maximum temperature, the Ar gas injection was switched to a 1% H_2 –Ar gas, and the reduction was allowed to proceed for various durations. The reduced powder was cooled to 100 °C in an Ar gas flow, and removed from the furnace at room temperature.

The powder samples were blended with MIBK and a trace high-molecular-weight polymer dispersant at a powder-to-MIBK weight ratio of 1:50, and they were stirred in a paint shaker with 0.3 mm ϕ zirconia beads to obtain a homogeneous dispersion. The particles were polished to a crystallite size of 23–35 nm by properly adjusting the polishing time that was monitored using the XRD Scherrer method. Details of the dispersion preparation method have been provided in ref. 25.

II-3 Characterization and calculation methods

A series of powder samples was subjected to powder-XRD measurement using Cu-K α radiation with an X'Pert-PRO/MPD apparatus (Spectris Co., Ltd., Tokyo, Japan). The diffraction angle was calibrated with a Si standard (NIST640e).

The nanoparticulate dispersions of the reduced products were further diluted with MIBK to a filler concentration of 0.05 wt%, injected into a transparent cell with a 1 mm optical path length, and its transmittance was measured with a U-4100 spectrophotometer (Hitachi High-Tech Corporation, Tokyo, Japan). The molar absorption coefficient was determined from the transmittance measurements. Visible light transmittance (VLT) was measured according to code JIS R3106. We measured solar transmittance according to JIS R3106 up to a wavelength of 2100 nm (ST21). It is defined by

$$\sum_{\lambda=300\;\mathrm{nm}}^{2100\;\mathrm{nm}} S(\lambda) T(\lambda) \Delta \lambda \left/ \sum_{\lambda=300\;\mathrm{nm}}^{2100\;\mathrm{nm}} S(\lambda) \Delta \lambda, \right.$$

where $S(\lambda)$ is the spectral distribution of the solar radiation, $T(\lambda)$ is the spectral transmittance, and $\Delta\lambda$ is an increment of wavelength. Hunter's chromatic indices, L^* , a^* , and b^* as well as RGB color indices were calculated according to JIS Z8701 and JIS Z8729.

Dispersions of neutral-toned ${\rm In_2O_3:Sn}$ (ENAM Optoelectronic Material Co., Ltd., Funan, China; hereafter referred to as ITO) and bluish fully-reduced Cs-HTB prepared by the GFR process¹⁹ (hereafter referred to as CWOTM) were prepared as color-tone references.

Powder size and morphology in those dispersions were observed using a transmission electron microscope (TEM) (HF-2200, Hitachi High-Tech Corp., Tokyo, Japan) operated at an acceleration voltage of 200 keV.

X-ray photoelectron spectroscopy (XPS) was performed using a Versa Probe II instrument (Ulvac-phi Inc., Chigasaki, Japan). As ion cleaning is known to reduce the sample surface, ¹⁹ the measurements were performed on freshly as-produced powders. Okada *et al.* ¹⁹ confirmed that Cs-HTB powders qualitatively give the same results as the bulk vacuum-cleaved surface. The precise position of the Fermi energy, $E_{\rm F}$, in the bottom of the conduction band was calibrated by evaporating Au on the second measurement.

First-principles calculations were performed in the Vienna *ab initio* simulation package (VASP),^{26–29} a calculation software based on plane-wave basis. Density functional theory with DFT+U method, which was confirmed to reproduce the optical properties of Cs_{0.33}WO₃ with high accuracy, was used.¹⁸ The +U values of the d orbital of W and the p orbital of O were 3.8 eV and 8.9 eV, respectively, as obtained in previous research.¹⁸

The optical properties of a material are defined by dielectric functions. Here, we applied the calculation method described in ref. 18. The dielectric function ε comprises a real part ε_1 and an imaginary part ε_2 , as shown in eqn (1). The two parts are connected by the Kramers–Krönig relation shown in eqn (2). Briefly, if the imaginary part of the dielectric function can be accurately obtained up to the high-energy region, the whole

Paper RSC Advances

dielectric function can be obtained by the Kramers-Krönig relation.

$$\varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega), \tag{1}$$

$$\varepsilon_1(\omega) = 1 + \frac{2}{\pi} P \int_0^\infty \frac{\varepsilon_2(\omega')\omega'}{\omega'^2 - \omega^2} d\omega'. \tag{2}$$

 ε_2 corresponds to the absorption of light energy by electrons, and comprises the Lorentz term (the contribution of interband transitions given by eqn (3)), and the Drude term (the contribution of free electrons). The former term, which is a 3 \times 3 Cartesian tensor, is given by

$$\varepsilon_{2,\alpha\beta}^{\text{Lorentz}}(\omega) = \frac{4\pi^2 e^2}{V} \lim_{q \to 0} \frac{1}{q^2} \sum_{c,v,k} 2w_k \delta(E_{c,k+q} - E_{v,k+q} - \omega) \\
\times \langle u_{c,k+qe_\alpha} | u_{v,k} \rangle \langle u_{c,k+qe_\beta} | u_{v,k} \rangle^*, \tag{3}$$

where V is the volume of the unit cell, and c and v are suffixes indicating the valence band and conduction band, respectively. E is the energy level, e is the unit vector with subscripts α and β denoting Cartesian coordinates, u is the periodic part of the orbit, and w_k is the weight of each set of k points.³⁰

Cs-HTB is a conductor whose Fermi energy crosses the conduction band. Therefore, both the Lorentz and Drude terms are necessary. The Drude terms are functions of the plasma frequency ω_p (given by eqn (4)) and the relaxation time $\tau=\frac{1}{\gamma}$. They are expressed by eqn (5) and (6):

$$\omega_{\rm p} = \frac{ne^2}{\varepsilon_0 m^*} = \frac{4\pi e^2}{V} \sum f_{nk} \frac{\partial^2 E}{\partial k \partial k},\tag{4}$$

$$\varepsilon_1^{\text{drude}}(\omega) = 1 - \frac{\omega_p^2}{\omega^2 + \gamma^2},$$
(5)

$$\varepsilon_2^{\text{drude}}(\omega) = \frac{\omega_p^2 \gamma}{\omega(\omega^2 + \gamma^2)}.$$
(6)

In eqn (4), f_{nk} is the Fermi-Dirac distribution function, n is the free electron density, e is the electron charge, ε_0 is the dielectric constant in a vacuum, and m^* is the effective mass of an electron.

Eqn (4) computes the plasma frequency from the band structure. The relaxation time indicates the effect of scattering caused by electron–phonon, electron–electron, and electron–lattice defect interactions. However, as γ does not appreciably affect the dielectric function, $\hbar \gamma$ was set to 0.1 eV in this study.

III Results and discussion

III-1 Optical properties of Cs-polytungstate

 $\mathrm{Cs_4W_{11}O_{35}}$ is a slightly greenish, almost white powder. Its MIBK dispersion is also white. Immediately upon reduction at 700–900 °C, a blue coloration occurred. As the reduction progressed, the color changed from sky blue to deeper blue to dark blue. Fig. 2(a) shows the color changes of the MIBK-dispersed CPT during $\mathrm{Cs_4W_{11}O_{35}}$ reduction at high temperature, and Fig. 2(b)

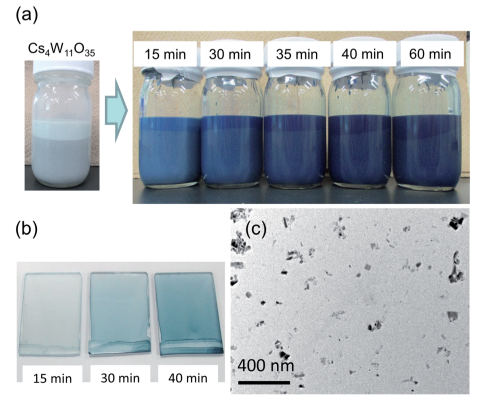


Fig. 2 Color changes of (a) dispersed liquids in settled beads and (b) flow-coated glass plates of CPTs obtained by reducing $Cs_4W_{11}O_{35}$ at 800 °C for various durations in a 1% H_2-N_2 gas flow. (c) TEM image of CPT dispersion reduced for 30 min.

shows their flow-coated films on a soda-lime glass. At reduction times of 15 minutes or less, the luminous color tone resembled that of ITO; after 40 minutes, it resembled that of CWOTM. The bluish tone of the flow coating (using 0.05 wt% CPT dispersion) increased as the b^* value decreased from 4.83 to -2.96 and then to -6.17. The TEM micrograph in Fig. 2(c) corresponds to the dispersed CPT nanoparticles reduced for 30 minutes at 800 °C and shows a typical image of CPT dispersions. The crystal shape is irregular because of the milling process, and the size scattered ranges from a few nm to 100 nm. The XRD Scherrer crystallite size was measured as 23–35 nm for every dispersion.

Panels (a)–(c) of Fig. 3 plot the molar absorption coefficients of the CPT dispersions as functions of energy. Strong double-peaked absorptions appear in the 0.6–1.8 eV range. The low-energy (low-*E*) and high-energy (high-*E*) peaks are attributable to LSPR and polaronic absorptions, respectively, and the hidden intermediate LSPR peak arises from the crystal anisotropy.²⁰ These absorptions strengthened with increasing reduction

temperature and/or time. The absorption obtained by heating at 800 $^{\circ}$ C for 15 minutes was attained by heating at 700 $^{\circ}$ C for 1 hour. No absorption occurred by heating at 550 $^{\circ}$ C for 1 hour. Conversely, the state of 30 minutes at 800 $^{\circ}$ C was reached after only 10 minutes at 900 $^{\circ}$ C. Therefore, the following analysis concentrates on the treatment at 800 $^{\circ}$ C.

Fig. 3(c) is an enlarged view of the visible-range absorption curves in Fig. 3(a). The profile of $Cs_4W_{11}O_{35}$ monotonously decreased from blue to red. At the blue side of the spectrum, the absorption decreased with increasing reduction time. Conversely, at the red side of the spectrum, the absorption remarkably increased with increasing LSPR absorption over time. Fig. 3(d) shows the series of transmission profiles of the samples with various CPT concentrations in dispersion, reduced for 35 minutes at 800 °C. The curves were derived from the molar absorption coefficient and the Lambert–Beer equation. Relative to the profile of CWO^{TM} (fully-reduced Cs-HTB produced by the normal route of GFR synthesis 19), the CPT

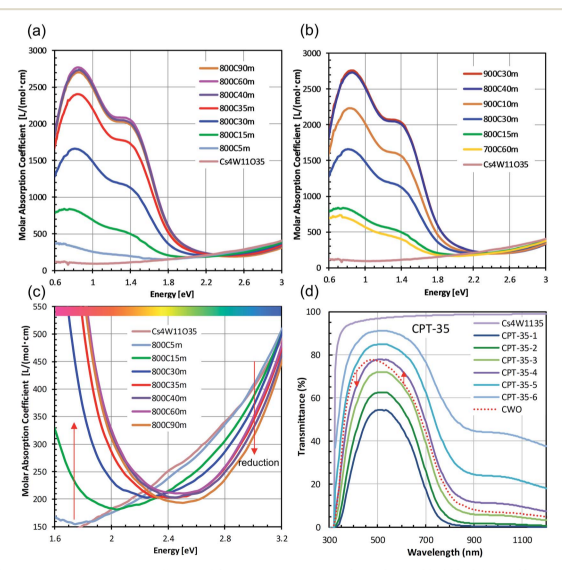


Fig. 3 (a-c) Measured molar absorption coefficients of CPT obtained by reducing $Cs_4W_{11}O_{35}$ at 700-900 °C for 15-60 min, and (d) transmittance profiles of the 800 °C - 35 min specimen, assuming Lambert-Beer's law.

Paper **RSC Advances**

transmission was reduced in the blue regions and increased in the red regions. Accordingly, the CPT series exhibits a more tintneutralized profile than CWOTM.

Here, it is noted that the optical profiles of the CPT dispersions shown above are highly consistent in terms of the degree of reduction even though certain irregularities in the size and morphology of the nanoparticles are observed in TEM images. This trend arises from the ensemble inhomogeneity effect of nanoparticle dispersions^{20,25} that generally broadens the absorption band and red-shifts the absorption peak, thereby masking the variation in individual Mie scattered waves due to differences in shape and physical properties.

Fig. 4 shows the changes in transmittance profiles, solar transmittance, CIE color indices, and RGB color indices of the dispersions reduced at 800 °C. For comparison, the evaluation results of neutral-colored ITO dispersion (manufactured by ENAM) and blue-tinted CWOTM dispersion synthesized by the normal route are also shown. As these optical indices depend on the transmittance value, the visible light transmittance (VLT) in

this comparison was fixed at 72.3%. The corresponding L^* range was 88.12 \pm 0.22.

The reduction of Cs₄W₁₁O₃₅ was accompanied by a large NIR absorption around 1500 nm (Fig. 4(a)), almost equaling that of CWO™ after 60 minutes' reduction. Owing to polaronic excitation, the absorption around 850 nm was characteristically stronger in CPT than in ITO. In the visible region (Fig. 4(b)), the transmittance of CPT in the blue region (relative to that of ITO) gradually increased with reduction time, approaching that of CWO™ at 60 min. In the red region, the transmittance of CPT sharply decreased owing to neighboring NIR absorption. The transmittance in this region was comparable to that of ITO at 15 min, and approached that of CWO™ at 60 min.

The NIR shielding effect increased with reduction time. After 20 minutes of reduction, the solar transmittance (ST21) was lower in CPT than in ITO (Fig. 4(c)); after 50 minutes, the transmittances in CPT and CWOTM were equal. Similarly, the CPT transmittance in the near-infrared region (T900, $\lambda = 900$ nm) was below that of ITO at 15 minutes and reached the CWOTM level at

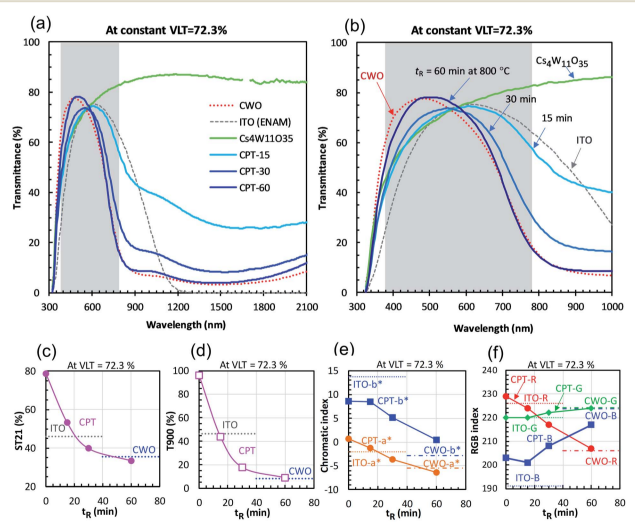


Fig. 4 (a and b) Transmittance profiles of CPT and (for comparison) fully-reduced CWO and ITO at constant visible light transmittance (VLT = 72.3%). The profile of $Cs_4W_{11}O_{35}$ at VLT = 72.3% is also shown. Reduction time dependence of (c) solar transmittance, (d) transmittance at 900 nm, (e) CIE color indices, and (f) RGB indices.

60 min (Fig. 4(d)). That is, the solar shielding property was higher in CPT than in ITO after 20 minutes of reduction. This high shielding is attributable to high absorption of CPT in the 700–1000 nm region, where the solar intensity is high.

Fig. 4(e) shows the changes in the CIE color indices (a^* and b^*) as functions of reducing time. As the reducing time increased, a^* changed from magenta to green while b^* changed steeply from yellow to blue. Even after 60 minutes of reduction, the b^* of CPT remained positive, indicating a weaker bluish tone than in CWOTM (with $b^* = -2.89$). Similarly, the R (red) index of CPT in the RGB color system linearly decreased with reduction time, whereas the B (blue) index increased with reduction time after a brief decline at 15 minutes. In summary, although the bluish tint in white $Cs_4W_{11}O_{35}$ generally strengthened with reduction time, a less-bluish color tone was compatible with the high solar shielding effect of CWOTM in the 30-60 min reduction time at 800 °C.

III-2 Structural changes during Cs₄W₁₁O₃₅ reduction

The structural factors responsible for the optical characteristics in Fig. 2–4 are presented. Fig. 5 shows the changing XRD powder patterns of $Cs_4W_{11}O_{35}$ reduced at 800 °C. The asproduced $Cs_4W_{11}O_{35}$ exhibited almost a single-phase pattern, consistent with an orthorhombic phase of space group $Pbc2_1$ as reported by Solodovnikov $et\ al.^{31}$ As the reduction time increased, the reflections of the orthorhombic $Cs_4W_{11}O_{35}$ phase gradually decreased (almost disappearing at 40 minutes), while

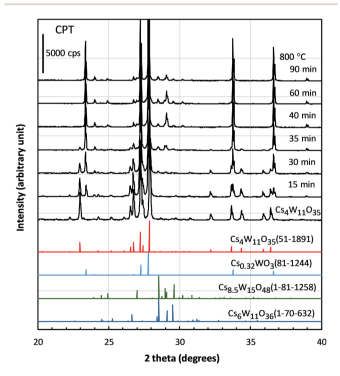


Fig. 5 XRD profiles of $Cs_4W_{11}O_{35}$ and the specimens reduced at 800 °C for different durations. Also shown are the profiles of $Cs_4W_{11}O_{35}$ (51-1891), $Cs_{0.32}WO_3$ (81-1244), $Cs_{8.5}W_{15}O_{48}$ (01-81-1258), and $Cs_6W_{11}O_{36}$ (01-70-632) extracted from the International Centre for Diffraction Data.

the reflections of the hexagonal $Cs_{0.32}WO_3$ phase³² gradually increased and dominated after 30 minutes. The space group of the hexagonal phase is $P6_3/mcm$ as originally determined by Magnéli³³ and as reported in recent works.^{19,32,34} The excess peaks around $2\theta = 28-31^{\circ}$ that appeared in the later stage of reduction matched only $Cs_{8.5}W_{15}O_{48}$ (space group: $R\bar{3}m$) or $Cs_6W_{11}O_{36}$ (space group: Aa) among the ICDD data base. Although the intensity distributions do not completely match those of Cs-rich $Cs_{8.5}W_{15}O_{48}$ or $Cs_6W_{11}O_{36}$, speculation exists that some modifications of these phases precipitated because the Cs composition becomes excessive with the growth of Wrich $Cs_{0.32}WO_3 = Cs_4W_{12}O_{36}$ phase. From the initial stage to 40 minutes of reduction, the powder-XRD characterization revealed a mixed-phase CPT comprising $Cs_4W_{11}O_{35}$ and $Cs_{0.32}WO_3$.

The lattice constants of the hexagonal components were obtained by the Rietveld method assuming $P6_3/mcm$. The lattice constants of the orthorhombic phase were determined by the Pawley method assuming $Pbc2_1$ because the precise atomic positions were undetermined. Fig. 6 shows the changing lattice constants with increasing reduction time in the hexagonal a-c space. The $Cs_4W_{11}O_{35}$ lattice is classified as orthorhombic, but in the model of Solodovnikov $et\ al.$, 31 it can also be regarded as a modified hexagonal lattice with W- and O-deficient planes periodically inserted in $\{020\}_{ORTH}//\{100\}_{HEX}$. Here the orthorhombic lattice constants were converted to hexagonal lattice constants by the formula: 31

$$a_{\text{ORTH}} = 2a_{\text{HEX}}, \ b_{\text{ORTH}} = 4\sqrt{3}a_{\text{HEX}}, \ \text{and} \ c_{\text{ORTH}} = c_{\text{HEX}}.$$
 (7)

As the reduction proceeded, the CPT was decomposed into orthorhombic and hexagonal fractions. The hexagonal

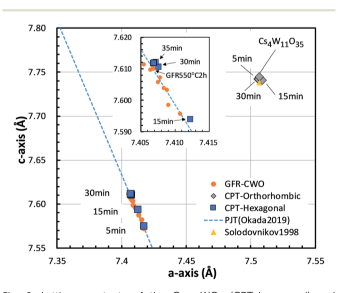


Fig. 6 Lattice constants of the $Cs_{0.32}WO_3$ (CPT-hexagonal) and $Cs_4W_{11}O_{35}$ (CPT-orthorhombic) portions in CPTs formed by reduction at 800 °C for various times. The c-axis versus a-axis is plotted in the hexagonal basis. The plot is compared with those of $Cs_4W_{11}O_{35}$ by Solodovnikov et al. ³¹ and $Cs_{0.32}WO_{3-y}$ produced by gas flow reduction (GFR-CWO) at 550 °C for varying reduction times. ¹⁹ The dashed line shows the pseudo Jahn–Teller-induced variation by Okada et al. ¹⁹

Paper RSC Advances

components were separated at positions of smaller a and c than the orthorhombic component. At later reduction times, the hexagonal a-axis and c-axis contracted and elongated, respectively, shifting to positions close to those of fully-reduced CWOTM, where they saturated. In recent experiments, 17,19 we showed that as the Cs and/or Vo concentration increases, the lattice constants of GFR-derived CWO™ move along the dashed line in Fig. 6 to the upper left. This behavior is triggered by electron donation to W-5d in the conduction band, which destabilizes the pseudo Jahn-Teller (PJT) distortion. 17,35 The lattice constants of the GFR-derived CWOTM were constant after two hours' reduction at 550 °C around the point labeled 'GFR550°C2h' in Fig. 6. When the reduction time exceeded four hours at 550 °C, WO₂ and W began precipitating. In the present reduction of Cs₄W₁₁O₃₅, the lattice constants similarly converged toward the full reduction saturation point as the reduction proceeded. This behavior can reasonably be attributed to destabilization of the PJT distortion as proposed in Okada et al.19 Clearly, the reduction process injects electrons into the W-5d orbitals of the conduction band.

III-3 Electronic changes during Cs₄W₁₁O₃₅ reduction

This subsection examines the factors in electronic structures that explain the optical properties shown in Fig. 2-4. Fig. 7 shows the changes in the W-4f XPS spectrum with increasing reduction time. As the reduction proceeded, the $W_{7/2}^{5+}$ peak at around 33.5 eV increased while the $W_{7/2}^{6+}$ peak at around 35.2 eV decreased. When plotted as a function of reduction time, the peak area ratio of W⁵⁺/W⁶⁺ increased to its maximum at 40 minutes, then decreased at 60 minutes. Similar changes have been observed in GFR-derived samples.19 Okada et al. determined the amount of Vos by chemical analysis, and concluded that Vos produces W5+ that supplies electrons to the conduction band, thus destabilizing the PIT distortion. Machida et al.20 analytically decomposed the optical absorption peaks into three components: a polaron peak and two anisotropic plasmon peaks. After extrapolating the intensity change of the polaronic component to zero reduction time, they found that the polaron component vanishes in the absence of V_O. Furthermore, in first-principles calculations, Yoshio and Ada chi^{18} showed that the 1.6 eV peak of ϵ_2 is correlated with polaronic absorption and requires the presence of Vo. They also reported that when Vo is generated, charges are produced at neighboring W⁶⁺ to generate a set of W⁵⁺s. Therefore, the W⁵⁺ increase in Fig. 7 is attributable to the generation and increase of V_Os by the reduction of Cs₄W₁₁O₃₅.

Next, we observe the XPS valence band. As shown in Fig. 8(a), the reduction of Cs₄W₁₁O₃₅ decreased the broad O-2p peaked at 8 eV and 10 eV below the band gap. The weak peak at the bottom of the conduction band (labeled "A") grew as the reduction progressed. Peak A corresponds to the contribution of free and localized electrons^{17,18} existing in W-5d in the lower conduction band. In the non-conductor Cs₄W₁₁O₃₅, peak A was caused by a slight reduction at the sample surface under the intense Xrays, which may be disregarded. At reduction times exceeding 40 min, the lower conduction band should be fully populated

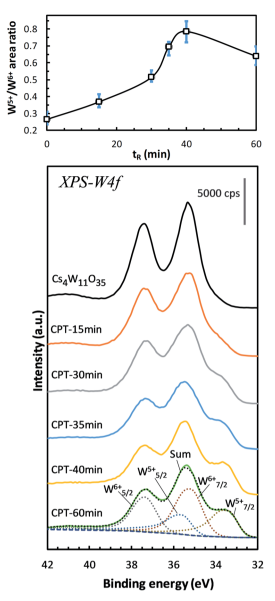


Fig. 7 W-4f spectra of Cs₄W₁₁O₃₅ and CPTs (15 $\leq t_{R} \leq$ 60 min), showing the increasing intensities of W⁵⁺ relative to those of W⁶⁺ versus reducing time $t_{\rm R}$. The growing W⁵⁺ indicates the production of oxygen vacancies.19

with free electrons, in analogous to fully-reduced CWOTM; indeed, the $E_{\rm F}$ crosses peak A at its half maximum. However, where the reduction was small, the peak height gradually decreased while simultaneously shifting toward the high- $E_{\rm B}$ side (valence-band side) with nearly zero density of states at $E_{\rm F}$. This phenomenon is considered to arise from the Coulomb gap.36 In the archetype tungsten bronze, NaxWO3, the decreasing electron density at $E_{\rm F}$ with decreasing Na content has been observed as the Coulomb gap. 37-39 As more electrons populate the conduction band, the Coulomb gap decreases. Because the Coulomb gap is caused by long-range interactions among trapped electrons, its presence in the CPT system further evidences electron-trapping in this system.

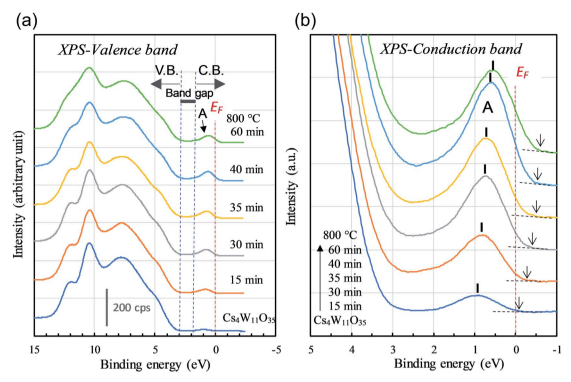


Fig. 8 XPS (a) valence- and (b) conduction-band spectra of $Cs_4W_{11}O_{35}$ and CPTs as the reducing time t_R varied from 15 to 60 min. The number of conduction-band electrons increases with t_R .

The above observations imply that when $\mathrm{Cs_4W_{11}O_{35}}$ is reduced at high temperatures, the electrons are donated to the conduction band by eliminating the W-defected planes caused by the orthorhombic-to-hexagonal transformation. Meanwhile, the parallel generation of $\mathrm{V_O}$ provides additional electrons to the conduction band. The electrons generated by both processes boost the NIR absorption capability.

Finally, the observed changes in electronic structures were examined by first-principles calculations.

Fig. 9 shows the crystal structures of (a) $Cs_4W_{11}O_{35}$ and (b) $Cs_4W_{12}O_{36}$. The structure in Fig. 9(b) is equivalent to $Cs_{0.33}WO_3$ reoriented in an orthorhombic basis for structural comparison with $Cs_4W_{11}O_{35}$. The structure in Fig. 9(a) is that of Fig. 9(b) with the removal of W and O on the (010) plane. The first-principles calculations considered the terminal structures of Fig. 9(a) and (b), and the W-deficient and O-deficient versions of the structure in Fig. 9(b).

As shown in Fig. 10, the conduction and valence bands of $Cs_4W_{11}O_{35}$ and $Cs_4W_{12}O_{36}$ have similar overall dispersion, although the position of E_F is different. The E_F locates in the band gap of the former, and in the lower conduction band of the latter. Therefore, $Cs_4W_{11}O_{35}$ is a non-conductor and $Cs_4W_{12}O_{36}$ is a conductor. With fully-arranged W and O atoms and a complete hexagonal WO_3 network, $Cs_4W_{12}O_{36}$ can inject its Cs^+ electrons into the W-5d orbitals, conferring a conducting property. 6,17,18

In the $Cs_6W_{17}O_{54}$ of Fig. 10(c), one W has been removed from $Cs_6W_{18}O_{54}$, the cell of which is 1.5 times larger along the *b*-axis direction of $Cs_4W_{12}O_{36}$. Although the charge is formally neutral in the chemical formula $Cs_6W_{17}O_{54} = 3Cs_2O \cdot 17WO_3$, the

material is considered as a conductor because $E_{\rm F}$ crosses the lower conduction band. When the W defect (V_W) was created, many localized levels were generated and the band gap was altered, implying an intrinsic change in the band structure. In contrast, when Cs is varied in Cs_xWO₃ (not shown here), the band structure is almost unchanged (only the $E_{\rm F}$ position changes).¹⁸

In the structure of Fig. 10(d), one O has been removed from the Cs–O plane of $Cs_4W_{12}O_{36}$. Additional electron orbitals generated in the band gap were partially localized in specific directions. Essentially, the same result was reported in a more

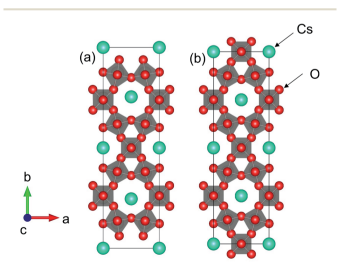


Fig. 9 Orthorhombic (001) projections of (a) $Cs_4W_{11}O_{35}$ and (b) $Cs_4W_{12}O_{36}$ ($Cs_{0.33}WO_3$).

Paper RSC Advances

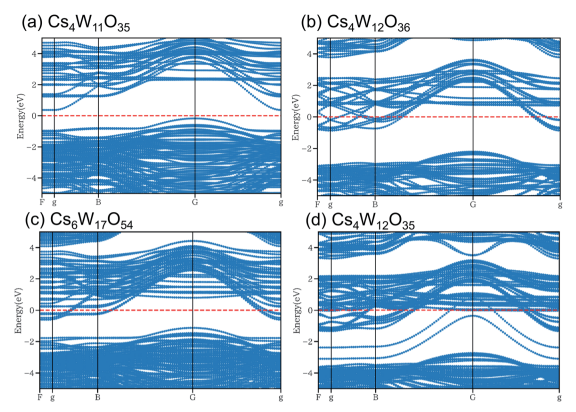


Fig. 10 Band structures of (a) $Cs_4W_{12}O_{35}$, (b) $Cs_4W_{12}O_{36}$ = $Cs_{0.33}WO_3$, (c) $Cs_6W_{17}O_{54}$ and (d) $Cs_4W_{12}O_{35}$ calculated by DFT+U method. Red dashed lines denote the Fermi energy.

detailed calculation ¹⁸ that included $V_{\rm O}s$ residing at multiple sites in both the Cs–O and W–O planes. When $V_{\rm O}$ is generated, the band structure changes as observed in $V_{\rm W}$.

According to the XRD observations, the reduction process induced a structural transformation from $Cs_4W_{11}O_{35}$ to $Cs_4W_{12}O_{36}$. The starting $Cs_4W_{11}O_{35}$ contained fewer W and O atoms than $Cs_4W_{12}O_{36}$. In part, these W and O deficiencies are probably inherited by the hexagonal CPT. As the reduction proceeded, the amounts of V_W and V_O decreased and increased, respectively. The V_W decrease corresponded to the changing band structures in Fig. $10(a) \rightarrow (c) \rightarrow (b)$, whereas the V_O increase corresponded to the changes in Fig. $10(a) \rightarrow (b) \rightarrow (d)$. In both cases, the E_F was displaced into the conduction band and the number of conduction electrons increased. Therefore, peak A in the XPS spectrum grew with increasing reduction time (Fig. 8), and the PJT destabilization shifted the lattice constants (Fig. 6).

Based on these band structures, the dielectric functions (including the Drude term) were calculated, and their real and imaginary parts are plotted in panels (a) and (b) of Fig. 11, respectively. At $\epsilon_1=0$ in Fig. 11(a), the screened plasma frequency (\varOmega_{sp}) is seen to increase in the order of $Cs_4W_{11}O_{35} \rightarrow Cs_6W_{17}O_{54} \rightarrow Cs_4W_{12}O_{36} \rightarrow Cs_4W_{12}O_{35}$. The NIR absorption is expected to increase in the same order, although experimentally, the disappearance of V_W accompanies the generation of V_O .

As clarified in Fig. 11(b), the imaginary part ε_2 of CPT and CWOTM is generally small in the visible-light region, as expected. This result explains the high visible transparency of

these materials. At 3.3 eV in the blue region of the spectrum, the interband transition dictates the optical absorption. Thus, the $\mathrm{Cs_4W_{11}O_{35}}$ absorption is enlarged by the narrow band gap. On the other hand, at 1.6 eV in the red region, the $\mathrm{Cs_4W_{12}O_{35}}$ absorption is enlarged by the high Ω_{sp} . Therefore, the transmitted light in the red region should also decrease in the order of increasing Ω_{sp} as stated above.

Fig. 11(c) shows the Mie scattering efficiency of a 20 nm-sized spherical particle calculated using these dielectric functions. Comparing panels (b) and (c) of Fig. 11 with the experimentally observed curves of molar absorption coefficients (Fig. 3(c)), one finds that the dielectric functions well reproduce the optical change in the visible region induced by the reduction process. Notably, the V_W increased the light absorption at the high-E side. The V_W amount is expected to be lower in the GFR-method than in Wöhler's method, but the V_O amount should be comparable in both methods. The results in Fig. 11 show that V_W increased and decreased the absorption at the high- and low-E sides, respectively, causing a more neutral shift in color tone than in normal CWO^{TM} .

According to these calculations, the less-bluish CPT obtained in this study than in normal-route CWOTM was attributed to the narrow band gap of $Cs_4W_{11}O_{35}$. CPT containing V_W and V_O can be categorized as tungsten bronze based on its structural and electronic features, but its energy-band structure slightly differs from that of CWOTM. From an application perspective, the band gap of CPT and the amount of electrons injected into the conduction band can be controlled by adjusting the high-

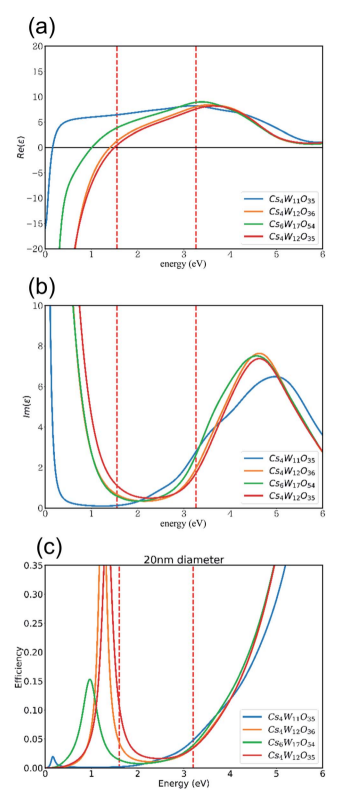


Fig. 11 (a) Real and (b) imaginary parts of the dielectric functions, and (c) the Mie scattering efficiency, of 20 nm-diameter particles of $Cs_4W_{11}O_{35}$, $Cs_4W_{12}O_{36}$ = $Cs_{0.33}WO_3$, $Cs_6W_{17}O_{54}$, and $Cs_4W_{12}O_{35}$.

temperature reduction condition of polytungstates, thereby achieving the desired NIR absorption effect with a low bluish color tint.

IV Conclusion

Revisiting Wöhler's method, we obtained Cs-doped tungsten bronze by reducing a Cs-polytungstate at high temperature. When orthorhombic $Cs_4W_{11}O_{35}$ was reduced at 700–900 °C, the NIR absorption and visual bluishness increased with reduction time. The structural phase transformation to hexagonal Cs_{0.32}-WO_{3-v} was observed to occur that accompanied the decrease in W defects. The linear change in the *c*-axis *versus a*-axis lattice constants of the hexagonal component was explained by destabilization of the pseudo Jahn-Teller distortion. Eventually, the lattice constants converged to those of fully-reduced CWOTM produced via the normal route. The XPS W⁵⁺ 4f signal increased as the reduction proceeded, confirming the formation of V_O. In the early reduction stages, the Coulomb gap was observed near the $E_{\rm F}$ of the Cs-polytung states, indicating the presence of trapped electrons. First-principles calculations confirmed that the decrease in W deficiency and/or the increase in O deficiency increased the $E_{\rm F}$ from the band gap to the conduction band. As Cs₄W₁₁O₃₅ has a comparatively narrow band gap and absorbs blue light, the reduced Cs-polytungstate decreased the transmission of blue wavelengths, achieving a less-bluish tone than fully-reduced CWOTM obtained by the normal route. A nonmetal-metal transition occurred when the Cs⁺-derived and Vo-derived electrons were donated to the W-5d orbitals in the conduction band, during the formation of the hexagonal lattice and the generation of Vos by the high-temperature reduction annealing.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

Experimental assistance provided by Ms. Kayo Yabuki on XPS measurements is greatly appreciated.

References

- 1 H. Takeda and K. Adachi, *J. Am. Ceram. Soc.*, 2007, **90**, 4059–4061.
- 2 Z. Yu, Y. Yao, J. Yao, L. Zhang, Z. Chen, Y. Gao and H. Luo, *J. Mater. Chem. A*, 2017, 5, 6019–6024.
- 3 L. Zheng, T. Xiong and K. W. Shah, *Sol. Energy*, 2019, **193**, 837–858.
- 4 W. Guo, C. Guo, N. Zheng, T. Sun and S. Liu, *Adv. Mater.*, 2017, **29**, 1604157.
- 5 C. Guo, S. Yin, H. Yu, S. Liu, Q. Dong, T. Goto, Z. Zhang, Y. Li and T. Sato, *Nanoscale*, 2013, 5, 6469.
- 6 J. B. Goodenough, Bull. Soc. Chim. Fr., 1965, 4, 1200.
- 7 O. F. Schirmer, V. Wittwer, G. Baur and G. Brandt, *J. Electrochem. Soc.*, 1977, **124**, 749.
- 8 E. Salje and B. Güttler, Philos. Mag. B, 1984, 50, 607-620.
- 9 A. Hussain, R. Gruehn and C. H. Rüscher, *J. Alloys Compd.*, 1997, **246**, 51–61.

Paper RSC Advances

- 10 G. A. Niklasson, J. Klasson and E. Olsson, *Electrochim. Acta*, 2001, **46**, 1967–1971.
- 11 K. Manthiram and A. P. Alivisatos, *J. Am. Chem. Soc.*, 2012, **134**, 3995–3998.
- 12 M. Kielwein, K. Saiki, G. Roth, J. Fink, G. Paasch and R. G. Egdell, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1995, 51, 10320–10335.
- 13 K. Adachi and T. Asahi, J. Mater. Res., 2012, 27, 965-970.
- 14 Y. Sato, M. Terauchi and K. Adachi, *J. Appl. Phys.*, 2012, **112**, 074308.
- 15 S. D. Lounis, E. L. Runnerstrom, A. Llordés and D. J. Milliron, *J. Phys. Chem. Lett.*, 2014, 5, 1564–1574.
- 16 L. Tegg, D. Cuskelly and V. J. Keast, *Plasmonics*, 2018, 13, 437–444.
- 17 S. Yoshio, M. Okada and K. Adachi, *J. Appl. Phys.*, 2018, **124**, 063109.
- 18 S. Yoshio and K. Adachi, Mater. Res. Express, 2019, 6, 026548.
- 19 M. Okada, K. Ono, S. Yoshio, H. Fukuyama and K. Adachi, *J. Am. Ceram. Soc.*, 2019, **102**, 5386–5400.
- 20 K. Machida, M. Okada and K. Adachi, J. Appl. Phys., 2019, 125, 103103.
- 21 S. Yoshio, K. Maki and K. Adachi, *J. Chem. Phys.*, 2016, 144, 234702.
- 22 A. Magnéli and B. Blomberg, *Acta Chem. Scand.*, 1951, 5, 372–378.
- 23 F. Wöhler, Ann. Phys. Phys. Chem., 1824, 78, 345-358.
- 24 E. O. Brimm, J. C. Brantley, J. H. Lorenz and M. H. Jellinek, *J. Am. Chem. Soc.*, 1951, 73, 5427–5432.
- 25 K. Machida and K. Adachi, *J. Phys. Chem. C*, 2016, **120**, 16919–16930.

- 26 G. Kresse and J. Furthmüller, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, 54, 11169–11186.
- 27 G. Kresse and D. Joubert, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **59**, 1758–1775.
- 28 G. Kresse and J. Hafner, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1993, **47**, 558–561.
- 29 G. Kresse and J. Furthmüller, *Comput. Mater. Sci.*, 1996, 6, 15–50.
- 30 M. Gajdoš, K. Hummer, G. Kresse, J. Furthmüller and F. Bechstedt, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2006, 73, 1–9.
- 31 S. F. Solodovnikov, N. V. Ivannikova, Z. A. Solodovnikova and E. S. Zolotova, *Inorg. Mater.*, 1998, **34**, 845.
- 32 L. Kihlborg and A. Hussain, *Mater. Res. Bull.*, 1979, **14**, 667–674
- 33 A. Magnéli, Acta Chem. Scand., 1953, 7, 315-324.
- 34 J. Oi, A. Kishimoto and T. Kudo, *J. Solid State Chem.*, 1993, **103**, 176–185.
- 35 I. B. Bersuker, Chem. Rev., 2013, 113, 1351-1390.
- 36 M. Lee, J. G. Massey, V. L. Nguyen and B. I. Shklovskii, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **60**, 1582–1591.
- 37 G. Hollinger, P. Pertosa, J. P. Doumerc, F. J. Himpsel and B. Reihl, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1985, 32, 1987–1991.
- 38 S. Raj, D. Hashimoto, H. Matsui, S. Souma, T. Sato, T. Takahashi, D. D. Sarma, P. Mahadevan and S. Oishi, *Phys. Rev. Lett.*, 2006, **96**, 3–6.
- 39 J. H. Davies and J. R. Franz, *Phys. Rev. Lett.*, 1986, 57, 475–478
- 40 Y. Sato, J. Appl. Phys., 2019, 126, 185107.