1 α-Ag₂WO₄ under microwave, electron beam and femtosecond laser

2 irradiations: Unveiling the relationship between morphology and

3 photoluminescence emissions

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22 ABSTRACT

23 In this study, the α-Ag₂WO₄ samples were successfully synthesized combined two methods, co-precipitation and microwave-assisted hydrothermal. Later, two different 24 irradiation processes: electron beam and femtosecond laser are applied. Unit cell changes 25 were shown by X-ray measurements and Rietveld analysis and compared with the results 26 27 obtained for first-principle calculations. The formation of oxygen vacancies on the surface of the particles after the irradiation process was revealed by XPS measurements. 28 Electron beam and femtosecond laser irradiations were found to cause expansion of the 29 unit cell, form oxygen vacancies on the surface, change the angle and distance between 30 31 O-Ag and O-W bonds, and modify the particle morphology to rod-, cube- and spherelike. The XANES measurements confirm that the local order of the W atoms is maintained 32 along the different irradiation processes. Based on the theoretical anlaysis of surfaces 33 34 investigation and Wulff construction, the contribution of (010) and (101) surfaces at the emission centers in 550 and 733 nm associated to the PL spectrum of α-Ag₂WO₄, was 35 established. 36

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40 *Keywords:* α-Ag₂WO₄; electron beam irradiation; femtosecond laser irradiation;
41 photoluminescence; morphology; surface band gap.

43 1. INTRODUCTION

Recently, alpha-silver tungstate (α -Ag₂WO₄) has attracted increasing attention from scientists due to its several unique characteristics. Among them, we can cite its nontoxicity [1–3], photoluminescence properties [4,5], and photocatalytic applications [1–6]. These properties can be improved according to the obtained morphologies by different synthesis methodologies, such as simple co-precipitation [6–8], conventional hydrothermal [9,10], or microwave-assisted hydrothermal (MAH) method [11].

50 When it comes to time-saving synthesis, the MAH method is considerably more 51 advantageous than other methodologies [12–15] since the use of microwave irradiation 52 triggers different processes between the forming clusters and the electromagnetic waves, 53 culminating in the formation of more oxygen defects and more structural/electronic 54 disorder effects in the structure of the material [16].

55 The properties derived from the interaction between electromagnetic waves and the material have been studied for several decades. It is known that the characterization 56 57 of structure or property responses of the material to these stimuli from the surrounding environment plays a key role in the understanding and rationalization of the structure-58 59 property relationship function in modern science and engineering [17,18]. In a recent study conducted by our group, α-Ag₂WO₄ was obtained by the MAH method. It was 60 observed that the photoluminescence of these samples migrated from red to blue, 61 according to the morphology and structural disorder presented [19]. 62

It is also known that factors such as temperature, time of synthesis, and presence of surfactants among others can alter the lattice structural properties of α -Ag₂WO₄ as well [6,20,21]. Regarding the influence of time of exposure to MAH, in a recent study performed by Laier et al. (2020) [20] both experimental and computational results revealed that at certain times of operation of the MAH system it was possible to obtain

samples with the highest active surface composition since they showed a higher density 68 of broken bonds and greater surface energy. The irradiation of electron beam and 69 femtosecond laser on the α -Ag₂WO₄ structure has attracted attention [10,11,22–32], 70 where the main focus is to elucidate the mechanism of growth of Ag nanoparticles on the 71 surface α–Ag₂WO₄ and the effects on the photoluminescence (PL) emissions [22, 32-35]. 72 In this work, we have tried to understand the phenomena provoked by the 73 interaction between α -Ag₂WO₄ and electron beam and femtosecond laser irradiation, 74 which were found to be able to promote changes in the lattice parameters, structure, and 75 morphology of the crystal. All of these changes alter the material properties, for example, 76 77 its electronic and magnetic properties, [22,36,37], PL emissions, and consequently its physical, chemical and/or biological applications [6,20,31,34,38]. Therefore, it is possible 78 to adjust the morphology, lattice, and electronic structures of α -Ag₂WO₄ materials 79 80 resulting from these modifications [19,39].

It should be noted that alterations in the lattice structure of the crystal may directly contribute to the variation in the number of vacancies. Once there is a critical correlation among lattice parameters, structural stability, electronic structure, band gap, and photoluminescence properties, it is important to have a broad understanding at the atomic-level of its bulk and surface.

In summary, This article came to increase knowledge about this structure and bring newness through the approach of explaining photoluminescence (PL), showing the surface that contributes to the property. It is well known that PL emission is the concentration of different defects: intrinsic (bulk and surface), extrinsic under interface, and structural order-disorder [32,40]. However, the surface contribution is still little known for the α -Ag₂WO₄ structure. For this purpose, we synthesized α -Ag₂WO₄ using the co-precipitation (CP) method, followed by the MAH method as a function of the

synthesis time (2, 4, 8, 16, and 32 min) and electron beam and femtosecond laser 93 94 irradiations. The crystals were characterized by different structural techniques, and their optical and PL properties were investigated. Theoretical calculations at the density 95 functional theory (DFT) level were performed to obtain atomic information of the 96 electronic structure of the material after irradiation. In addition, to understand the 97 relationship between PL emissions of the α -Ag₂WO₄ exposed surfaces, the corresponding 98 band gap was analyzed. The paper is organized as follows: section 2 describes the 99 experimental procedure (synthesis, characterization, and irradiation methods) and the 100 theoretical method, whereas section 3 shows the results and discussion on the structure, 101 102 morphology, and optical properties of α -Ag₂WO₄. Finally, we present our main conclusions in section 4. 103

104

105 2. EXPERIMENTAL SECTION

106 2.1. Synthesis of α-Ag₂WO₄ crystals

107 The α -Ag₂WO₄ samples were synthesized by the CP method, similar to the 108 description made by Foggi, *et al.* (2017) [41], being considered a volume of 30 mL of 109 ethanol, then kept under continuous stirring for 10 min; the as-obtained sample was 110 denoted as CP. The suspension followed by treatment in the MAH method (**Figure 1a,b**) 111 under the previous studies by [4,5,20,42], considering the treatment times of 2, 4, 8, 16, 112 and 32 min. These samples were denoted as MAH-2, MAH-4, MAH-8, MAH-16, and 113 MAH-32.

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115 **2.2. Irradiated samples (EI and FI)**

To obtain the irradiated samples, the set of MAH samples were submitted to two
different irradiations: electron beam irradiation (EI) (Figure 1c) and femtosecond laser

118	irradiation (FI) (Figure 1d). The EI process was conducted in a field emission scanning
119	electron microscope (Supra 35-VP; Carl Zeiss – Germany), using an acceleration voltage
120	of 30 kV for 2 minutes, the parameters were selected based on experiments reported in
121	the references [24,28,30]. The samples obtained through this process were denoted as EI-
122	2, EI-4, EI-8, EI-16, and EI-32. Regarding the femtosecond laser irradiation, the
123	procedure is according to Protocol I reported in reference [29], using a titanium/sapphire
124	laser (Femtopower Compact Pro, Femto Lasers) with pulses of 30 fs of full width at half
125	maximum, the wavelength of 800 nm, the repetition rate of 1 kHz and fluence of 60 J/cm ²
126	in diameter of the order of 20 μ m. The obtained samples were denoted as FI-2, FI-4, FI-
127	8, FI-16, and FI-32. Experimental characterizations of the samples are in Supplementary
128	Information.
129	
130	Figure 1.
131	

132 **2.3. Model systems and Theoretical Methods**

Density functional theory (DFT) calculations were performed with the 133 CRYSTAL17 program [43]. The B3LYP hybrid functional (doi:10.1063/1.464304) was 134 135 used for described the electron-electron interactions combined with the effective core pseudopotentials (ECP) derived by Apra [44] and Corà [45] which have been chosen for 136 described silver and tungsten, respectively, while the oxygen was described with the 8-137 411d11G basis set [46]. From the experimental results of Rietveld refinements of the CP 138 sample, we derived a theoretical reference model ($opt - \alpha$ -Ag₂WO₄) by full optimization 139 of the structural parameters of α -Ag₂WO₄. The accuracy in the evaluation of the Coulomb 140 and exchange was controlled by a set of tolerances with values of 10⁻⁸, 10⁻⁸, 10⁻⁸, 10⁻⁸, and 141 10^{-16} . A total mesh of 125 k-points was employed to sample uniformly the irreducible part 142

of the Brillouin zone. The vibrational modes at the Γ point were calculated by using the
numerical second derivates of the total energy. On the other hand, for the set of MAH,
EI, and FI samples, only the atomic positions were allow for relax. The relaxed structures
were used for the electronic structure calculations. Visualization of the unit cells was
performed using the VESTA program [47].
The electronic structure of the (010), (100), (001), (011), (101), and (110) surfaces

of α-Ag₂WO₄ were discussed from band gap energy values and the density of states
(DOS). Details computationals for the surfaces calculations can be found in previous
studies. [48]

152 The density broken bonds in the surfaces α -Ag₂WO₄ were calculated as

$$D_b = N_b/A,$$
 Eq. 1

where N_b is the number of broken bonds and A is the surface area. The number of broken bonds for each surface was taken from reference [21].

The polyhedron energy band gap energy value for the morphologies of α-Ag₂WO₄
was calculated as follows

158
$$E_{ap}^{polyhedron} = \sum_{i} C_{i} \times E_{gap}^{surf}$$
 Eq. 2

where $C_i = A^{surf} / A^{polyedron}$ is the ratio of the surface area (A^{surf}) to the total surface area of the polyhedron $(A^{polyedron})$ and E_{gap}^{surf} is the energy value of the corresponding surface.

162

163 **3. RESULTS AND DISCUSSION**

164 **3.1. Morphology**

165 α -Ag₂WO₄ samples obtained by the CP method using 70% of water and 30% of 166 ethanol, followed by treatment with MAH and exposure to EI and FI irradiations, were 167 investigated by FE-SEM, as shown in **Figure S1** and **Figure 2**. Through the CP method,

only hexagonal rod-like morphologies were observed, with preferential growth on the y-168 169 axis in the (010) direction (Figure S1). These characteristics were previously studied by our research group and are reported in reference [48]. In this method, the clusters of silver 170 and tungsten ions solvated with water and ethanol interact randomly in a polarization 171 process of complex clusters, depending on the moment of the permanent dipole of the 172 neighboring clusters. This short-range interaction induces the first links between the 173 174 crystal symmetry and its order/disorder. Then, to increase the symmetry and order of the crystal, there is a correlation between the rotation motions of permanent moments in the 175 different complex clusters of silver and tungstate to form the different surfaces. 176

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- 178

Figure 2.

179

180 Theoretical calculations reported in the literature show that the morphological shapes of α -Ag₂WO₄ are formed by the combination of the (010), (100), (001), (110), 181 (101), and (011) surfaces [21,48]. Although the theoretical results establish an ideal 182 parallelepiped-like morphology for α -Ag₂WO₄, characterized by the presence of (010), 183 (100), and (001) surfaces with predominant contributions of the (010) and (100) surfaces 184 185 and a minor proportion of the (001) surface, experimentally a hexagonal rod-like morphology was observed (Figure S1). Table 1 presents the values of E_{surf} according 186 to R. A. Roca and et al. [48], as well as the surface band gap energy (E_{gap}^{surf}) and the 187 percentage of contribution (% C_i) for each morphology of α -Ag₂WO₄. 188

189 The experimental rod-like morphology of α -Ag₂WO₄ for the CP sample 190 composed of the (010), (100), and (101) surfaces were obtained by destabilizing the E_{suf} 191 of the (010), (110), and (100) surfaces from 0.20 to 1.44 Jm^{-2} , 0.65 to 1.50 Jm^{-2} and 192 0.38 to 0.70 Jm^{-2} , respectively, as well as by stabilizing the (101) and (001) surfaces 193 from 0.68 to 0.23 Jm^{-2} and 0.53 to 0.22 Jm^{-2} , respectively (see Figure 3).

194

Figure 3.

195

In the MAH system, the monitoring of the experiment illustrated the time
dependence in relation to the morphological evolution of α-Ag₂WO₄ (Figures 2a-e).
Morphologies composed of cube-like (highlighted in orange) and rod-like (indicated in
blue) were observed in the samples treated with MAH at 2, 4, 8, and 16 min.
Theoretically, the cube-like morphology composed of the (010), (100), and (001) surfaces
can be obtained by stabilizing the (100) and (001) surfaces (Figure 3).

At 32 min of treatment with MAH, a process of redissolution and stabilization of 202 203 the (101) surface occurred, with concomitant destabilization of the (100) surface. This 204 information allows us to understand how the time variation of the MAH treatment in an 205 alcoholic environment influences particle growth. The microwave radiation induces a 206 structural disorder through the modification of the bond distance, consequently altering the internal rotation barrier and vibration of molecules. In turn, the stabilization of 207 previously unstable surfaces results in a new morphology [7,9]. Thus, at the highest 208 synthesis time (32 min) the MAH-32 sample (Figure 2e) presented no cubes – only rods 209 210 behaving similarly to those obtained in the CP sample (Figure S1). It is then believed 211 that at a longer exposure time to microwave irradiation, the cube-like morphology is 212 rearranged to a rod-like morphology.

213

Table 1. Calculated density broken bond (nm^{-2}) , surface energy (E_{surf}, Jm^{-2}) , surface band gap (E_{gap}^{surf}, eV) , percentage of contribution of each surface in the total area (% C_i) and polyhedron band gap energy for the morphologies of α -Ag₂WO₄.

Morphology	Surface	D_B	E _{surf}	% C _i	E_{gap}^{surf}	$E_{gap}^{polyhedron}$
Ideal	(010)	4.68	0.20	52.5	1.39	
$\alpha - Ag_2WO_4$	(100)	5.83	0.38	27.6	0.48	0.99
6-	(001)	6.04	0.53	19.8	0.65	
Rod-like	(010)		1.44	7.3	1.39	
$\alpha - Ag_2WO_4$	(001)		0.22	13.2	0.65	1.10
5	(101)	8.75	0.23	79.4	1.15	
Cube-like	(010)		0.20	42.9	1.39	
$\alpha - Ag_2WO_4$	(100)		0.30	28.6	0.48	0.92
c	(001)		0.30	28.6	0.65	

It can be inferred that the microwave treatment directly interferes with the crystal 218 morphology by either totally or partially dissolving the rod-liked crystals or altering the 219 220 internal rotation movements between the clusters. Thus, in the MAH method, the cubelike morphology is formed by the less energetic surfaces, i.e., (100) and (001), with a 221 222 decrease in disorder and an increase in symmetry (Figure 3). The same behavior was 223 observed in reference [28], in which the MAH method is employed to synthetize α -Ag₂WO₄ and different morphologies were obtained: hexagonal rod-like elongated, cubic-224 like, and triangular-like shapes. 225

Under electron beam irradiation, these morphologies change to a unique rod-like morphology (EI samples indicated by green color in Figures 2f-j), whereas under femtosecond laser irradiation they change to the mixed rod- (violet color) and sphere-like morphologies (yellow color) (FI samples in Figures 2k-o). The n-type semiconductor, α – Ag₂WO₄, when irradiated with electrons, grows on its surface metallic silver nanowires [10,11,30], thus forming silver vacancies inside the particle and causing a variation in the

surface energies of the crystalline structure. Therefore, the value of the (101) surface 232 drops from 0.68 to 0.23 Im^{-2} (Figure 3), becoming less energetic, while that of the (100) 233 surface jumps from 0.38 to 0.70 Im^{-2} , thus becoming more energetic. Even when varying 234 235 the synthesis times, all samples exhibit agglomerated nature with characteristics similar to those observed in the CP sample (Figure S1). In this micrograph, it can also be 236 observed that the hexagonal rod-like morphology is better defined, with few 237 238 imperfections. This fact can be considered an indication of the effect of electron beam irradiation on the morphology of α -Ag₂WO₄. 239

240 Under EI, the energy of the electron beam is transferred to the sample, causing different phenomena, such as atomic position change, surface distortions, electrostatic 241 charge, and local heating in the crystals. Depending on the voltage and the type of sample 242 243 electron beam will penetrate a few micrometers or nanometers deep into the sample. In 244 our case, the voltage is 30 keV, whose penetration is in the order of nanometers, causing surface defects and atomic mobility that culminate in the formation of Ag nanoparticles 245 246 on the surface of α -Ag₂WO₄ [11,49]. On the other hand, when a FI is applied, photons are absorbed by the sample to provoke an excitation in the crystal system. To return to 247 the ground state, the system segregates electrons, atoms, and ions from its structure, 248 forming a plasma plume ablation, which is highly energetic. This plume interacts with 249 250 the irradiated region, promoting energy exchanges, increasing temperature and pressure, 251 accelerating the speed of movement of the species, and ejecting particles. After this highly 252 energetic ablation process, the surface cools very quickly, which modify the surface structure of the material [37,50]. 253

Regarding the morphology obtained by the FI technique (**Figures 2k-o**), it is possible to observe in α -Ag₂WO₄ spherical particles with well-defined edges and irregular extensions, in addition to elongated and slightly curved rods. According to the

257	elementary mapping performed on the FI-8 sample (Figure 4), both the spheres and the
258	rods are composed of the same proportion of Ag, O, and W atoms, thus confirming the
259	change in particle morphology composed of the same α -Ag ₂ WO ₄ polymorph.
260	
261	Figure 4.
262	
263	The application of the FI technique leads to the formation of a spherical
264	morphology of α -Ag ₂ WO ₄ powders due to when the femtosecond laser beam interacts
265	with the α -Ag ₂ WO ₄ , a nonlinear and multiphoton ionization process occurs, which results
266	in the formation of a plasma plume above the irradiated zone [31,51]. The highly
267	energetic ablation process provokes the breaking of the bonds between the W-O and Ag-
268	O, atoms. Further, since the plasma itself can reach temperature and pressure values of
269	up to 1000 K and 10^{12} Pa, respectively [51], such extreme conditions can trigger the
270	sintering of the material that reminds in contact with the plasma Since the plasma plume,
271	which reaches values up to 1000 K, has a lifetime in the region of the ns [52], the plasma
272	can transfer heat to the material surrounding the irradiation zone. Thus, the material that
273	did not undergo sintering just experienced a melting process, resulting in the curvature of
274	its edges. This effect is considered to favor the coalescence of the previously observed
275	rods and cubes, giving rise to the morphology of the spheres.
276	The analysis in Figure 2 suggests that in addition to the morphological changes
277	already discussed, another relevant aspect is observed: the presence of silver
278	nanoparticles on the surface of samples as a result of electron beam and femtosecond laser
279	irradiations. This phenomenon has already been well discussed by our group
280	[10,11,24,31,38–40]. These changes in sample morphologies allow us to understand how

the surface structure can affect and modify the PL property of a semiconductor.

283

3.2. Unit cell and lattice parameters

284 The result of Rietveld refinement (Table S1) reveals that the lattice parameters and bond angle of all samples correspond to an orthorhombic structure and *Pn2n* spatial 285 group, which is in agreement with the crystallographic information file in ICSD 4165 286 card [53]. Moreover, the R_{Bragg} and chi² fitting parameters evidence the accordance 287 between the calculated data and the observed XRD patterns. The Rietveld refinement data 288 show the effect of irradiation on the parameters of the unit cell, which were found to 289 change. In general, when all MAH samples are irradiated by EI and FI, there is an 290 291 expansion in the cell volume, consequently altering the lattice of the samples.

292 **Figure 5** brings a comparison among the geometries for optimized ($opt-\alpha$ -AWO), neutral (CP), and irradiated structures (MAH-8, EI-8, and FI-8). From this figure, it is 293 294 possible to observe variations of the relative positions of the Ag, O, and W atoms on several different constituents $[WO_6]/[AgO_y]$ clusters (y = 7, 6, 4, and 2) of α -Ag₂WO₄. 295 296 Since the [AgO₂] and [AgO₄] clusters are found in the shell of the unit cell, changes are expected as a result of the strong interaction caused by irradiation, leading to nucleation 297 and formation of metallic Ag [24,54]. Thus, the comparison between the $opt-\alpha$ -AWO 298 299 geometries and MAH-8 structures for the sample treated with MAH indicates variations in the Ag–O distances and O–Ag–O angles, mainly in the [AgO₂] and [AgO₄] clusters. In 300 the [AgO₂] cluster, Ag–O increases from 2.246 to 2.461 Å, while O–Ag–O decreases 301 from 174.42 to 145.05 °. In the case of the Ag–O bond in [AgO₄] clusters, an increase in 302 two bond distances is observed. When the MAH-8 sample is exposed to EI, both Ag-O 303 304 distance and O-Ag-O angle in the [AgO₂] cluster remain almost unchanged, going from 2.461 to 2.455 Å and from 145.05 to 147.42 °, respectively, whereas the Ag–O distance 305 in the [AgO₄] clusters undergoes increases. On the other hand, when a femtosecond laser 306

is applied, the Ag–O distance in the [AgO₂] cluster decreases from 2.461 to 2.193 Å, while the O–Ag–O angle increases from 145.05 to 170.42 °. Regarding the [AgO₄] clusters, it is possible to note a shortening average of the Ag–O distance. With respect to the different types of [WO₆] clusters, in all cases, the W–O bond distances of the irradiated samples undergo variations corresponding to W1, in which an average lengthening is observed.

A careful analysis of the Ag–O and W–O bond distances for all samples were performed, and the observed values are listed in **Table S2**.

From Figure 5 and the values in Table S2, we can infer that the MAH treatment 315 316 induces a larger structural organization in the α -Ag₂WO₄, which is evidenced through the average lengthening of the Ag–O and W–O bonds in the $[WO_6]/[AgO_y]$ clusters (y = 7, 317 6, 4, and 2). Considering the EI, it can be noted that the irradiation with electrons leads 318 319 to structural changes in the [AgO₄] and [W1O₆] clusters. On the other hand, FI induces an average shortening of Ag–O and an average lengthening of W1–O bond distances in 320 321 the clusters of a-Ag₂WO₄ when compared with both MAH samples and the optimized 322 system. These results explain the constant changes in the crystal lattice and the size reduction of [AgO_y] clusters proposed by our group [22], which may also be induced by 323 324 rod-to-sphere morphology changes in the irradiated samples.

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- 326

Figure 5.

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To investigate the effect of irradiation on the particle surface, XPS measurements were conducted. Such analyses were performed in the samples that had their morphology and the PL property changed, that is, MAH-8, EI-8, and FI-8. Figure 6 shows the high-resolution spectra of O *1s*, which were fitted with three components. The strong peak around 530.03 eV, 529.83 eV, and 530.07 eV for the MAH-8, EI-8, and FI-8 samples, respectively, are attributed to the O atom in the α -Ag₂WO₄ lattice. The peak at 531.43 eV, 531.54 eV, and 531.92 eV, respectively, are assigned to oxygen defects in the lattice, such as oxygen vacancies (V_O) [55]. Lastly, the peak in the high binding energy, i.e., above 533 eV, is related to adsorbed oxygen on the surface [55,56].

Changes in the profile of the high-resolution spectrum of O Is indicate a high 338 percentage of adsorbed oxygen on the surface of the EI-8 and FI-8 samples in comparison 339 340 with MAH-8. Additionally, there is a decrease in the area percentage of the lattice oxygen to 69.97% for MAH-8, 48.00% for EI-8, and 37.66% for FI-8, as well as an increase in 341 342 the area percentage ascribed to defect oxygen to 20.98%, 27.88% and 38.10% for the 343 MAH, EI and FI samples, respectively. This indicates a possible formation of oxygen vacancies in the EI and FI samples. The vacancies formed are a charge compensation due 344 345 to destabilization of the (010) surface along the transformation of morphology from 346 cubes-like into rods-like.

Furthermore, surfaces can be stabilized by reducing surface charge density via different mechanisms: creating surface states, redistributing charge between atoms, removing atoms, adding charged impurities to the surface, or creating faces [57,58]. In the process of electron beam irradiation and femtosecond laser irradiations, we observed the destabilization of the surface (010), by the conversion of cubic particles into rods, which was compensated by the removal of surface oxygen atoms and by chemoadsorption of species (O^- and O^{2-}) on the surface, according to the XPS results.

The increment in oxygen vacancies and impurities, via FI, may be related to the fact that the experiment is carried out in an environment, which allows the exchange of charges and specie. It is known that the greater exposure of a given crystalline surface
influences the properties of materials [59]. An analyis of the theoretical results renders
that the (010) surfacepresent the highest value of surface energy in the rod morphology.
Extrapolating this data to literature [59], we suppose that to compensate the energetic
instability, superficial oxygen vacancies were created.

Figure S2a shows that the Ag *3d* spectra were better fitted using one component. The high-intensity peaks located at approximately 368 eV and 374 eV are related to the $3d_{5/2}$ and $3d_{3/2}$ orbitals, respectively, with Ag⁺ binding energy. No significant changes were observed among the spectra of the samples.

365 The XPS technique also provided information on the chemical environment of the W element, which is considered to be the lattice-forming atom. Figure S2b displays the 366 XPS spectra of the W 4f for the MAH-8, EI-8, and FI-8 samples. The binding energy 367 corresponds to the $4f_{7/2}$ and $4f_{5/2}$ orbitals of W⁶⁺ in the α -Ag₂WO₄ lattice [4,60,61]. The 368 XPS spectra reveal a similar profile in all samples, with no changes in the oxidation state 369 370 or chemical environment of the structural W atom, nor even for the EI and FI samples. 371 Therefore, it can be concluded that the irradiation process changes the environment around oxygen atoms due to the formation of metallic Ag and Vo vacancies, consequently 372 373 leading to distortions in the angle and bond distance of the Ag and W clusters (Table S2).

374

375

Figure 6.

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377 **3.3. Structural characterization**

Figure S3 displays the Raman spectra in the range of 60-960 cm⁻¹ at room temperature. According to Turkovič et al. (1977) [62], there are 21 known Raman modes for α -Ag₂WO₄ (6A_{1g}, 5A_{2g}, 5B_{1g}, and 5B_{2g}). As it can be observed in Figures S3a-d, six

Raman modes were detected for all samples (1A1g, 1A2g, 3B1g, and 1B2g); the remaining 381 382 modes were not detectable experimentally because of their low intensities. External vibrational modes of interaction between silver and tungstate ion and $[AgO_y]$ (y = 7, 6, 4, 383 2) clusters can be identified between 100 and 500 cm⁻¹, whereas internal vibrational 384 modes assigned to the vibrations in the [WO₆] clusters can be detected between 500 and 385 1000 cm⁻¹ [28,34,63]. It is possible to observe in both materials two intense bands: one at 386 102 cm⁻¹, attributed to the stretching of T'(Ag⁺/W⁶⁺) binding [64], and another at 877 cm⁻ 387 ¹, assigned to the symmetrical stretching of the W–O bond in the octahedral [WO₆] cluster 388 [28]. 389

The comparison between the relative experimental and theoretical positions of these vibrational modes is illustrated in **Figure S3e** and summarized in **Table S3**, confirming the good agreement between the experimental and theoretical modes.

Following the Raman discussion on the short-range structural disorder, XANES allowed us to visualize a local disorder, which in our case was around the tungsten atom. In the past decade, our research group has employed the XANES technique to study the local structure of various semiconducting metal oxides, such as tungstates, titanates, and vanadates among others [19,65–69].

398 Herein, we performed XANES measurements to investigate the effect of irradiation on the local structure around W atoms. Figure S4a displays the W-L1 XANES 399 spectra of the as-prepared CP sample and some standards used as reference samples 400 (Na₂WO₄, WO₂, and monoclinic WO₃). In Figure S4a, it is possible to observe a 401 pronounced peak (here denoted as P1) in the Na₂WO₄ spectrum, more specifically in the 402 pre-edge region, as well as a shoulder in the m-WO₃ standard and the CP sample spectra. 403 The physical origin of such electronic transition (peak P1) is described elsewhere 404 [19,70,71]. 405

It is well established that an intense pre-peak in the W-L1 edge XANES spectrum 406 407 has been found in materials constituted by tetrahedral units (WO₄), such as the Na₂WO₄ 408 compound [5,71]. The electronic transition responsible for P1 pre-peak is forbidden in 409 the materials that exhibit regular [WO₆] units. However, compounds presenting nonregular octahedral symmetry (distorted WO₆ clusters) have been found to present a less 410 411 intense pre-peak similar to that in the WO₃ spectrum [70,71]. From the analysis of the 412 spectrum of the CP sample, it is possible to observe its similarity with the WO₃ standard spectrum. It is then plausible to affirm the presence of distorted [WO₆] clusters in the as-413 prepared CP sample. Furthermore, by comparing the spectra of the α-Ag₂WO₄ samples 414 415 (Figure S4b) it is evident the similarity with the XANES spectra (both in the pre- and post-edge regions). Such behavior reveals that the microwave-assisted hydrothermal 416 417 treatment, as well as the irradiation processes (EI and FI), resulted in samples constituted 418 by distorted octahedral WO₆ clusters.

419

420 **3.4. Density of states**

The experimental optical band gap was estimated using the Wood-Tauc relation 421 422 [72,73] and the Kubelka-Munk equation [59,60], evidencing a slight variation from 3.09 423 to 3.32 eV (Figure S5) for the MAH, EI, and FI samples. The experimental results show that when the CP sample is submiiited to MAH treatment for 2 minutes, the band gap 424 value decreases from 3.13 eV to 3.08 eV. In contrast, for the MAH-4, MAH-8, MAH-16, 425 and MAH-32 samples, the band gap increases. Later, when these samples are under EI, 426 is observed that except for the EI-2 sample, the band gap decrease in all cases. For the FI-427 2 and FI-4 samples present an increase in the band gap, while for the FI-8, FI-16, and FI-428 32 samples, an opposite effect can be sensed. Structural order-disorder effects induced by 429

the radiation as well as the morphology, time process and shape powder can explain the
band gap decrease/increase of α-Ag₂WO₄.

The electronic structure of α-Ag₂WO₄ was analyzed from the partial density of 432 states (PDOS) of the 4d, 5d, and 2p orbitals of Ag, W, and O atoms, respectively (Figure 433 7), where the valence band maximum (VBM) is set to zero. In terms of contributing states, 434 the representation is similar for all samples, according to the composition of the valence 435 436 band (VB) and conduction band (CB) of the different samples. As already known, for α-Ag₂WO₄ the VB is formed by the hybridization of $4d_{xy}$ and O $2p_z$ orbitals, whereas the 437 bottom of the CB is mainly composed of W 5d orbitals, more specifically $5d_z^2$ orbitals. 438 439 However, differences in topology occur among the PDOS of neutral and irradiated samples. In the PDOS of the neutral sample, there is discontinuation in the VB, leading 440 to two distinct Ag 4d blocks. Such characteristic is maintained in the EI-8 sample, while 441 442 in the PDOS of the MAH-8 and FI-8 samples the double-peak structure disappears.

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- 444

Figure 7.

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In the MAH-8 and FI-8 samples (Figure 7c), localized states of the O $2p_x$, $2p_y$ and 446 $2p_z$ and Ag $4d_{xy}$ orbitals are evidenced in the Fermi region, as well as localized $5d_x^2 y^2$ and 447 $5d_z^2$ orbitals forming the bottom of the CB in these samples. The localized density of 2p448 and W 5d states situated at the Fermi level and the bottom of the CB, respectively, favors 449 450 the electronic transfer from the O 2p to W 5d states, which are necessary for the excitation process of the tungstate groups. These observations in the electronic properties of the 451 different samples are associated with the structural disorder induced by the irradiation in 452 the α -Ag₂WO₄ lattice, as seen in Figure 5. 453

Figure 8 shows the PL spectra of α -Ag ₂ WO ₄ samples measured at room
temperature and excited at 355 nm. The PL spectra encompass the whole visible region
and part of the near-infrared region, presenting a broadband profile. The small changes in
the profile of the PL spectrum between the same group are related to different degrees of
order/disorder in the O-Ag and O-W bond lengths. Therefore, by analyzing the PL
spectra we can understand how the MAH treatment time, and electron beam and
femtosecond laser irradiations affect the emission spectrum of α-Ag ₂ WO ₄ .
Figure 8.
The CP and EI samples with rod-like morphology (Figure S2 and Figure 2f-j)
have two maximum emission centers: around the green (550 nm) and the red (733 nm)
regions (Figure 8a,b). The FI sample with rod- and sphere-like morphologies presented
the same emission centers (Figure 2k-o and Figure 8c). When the sample is submitted
the same emission centers (Figure 2k-o and Figure 8c). When the sample is submitted to MAH at synthesis times between 2 and 16 min, two types of morphologies are observed
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478 The blue band is attributed to the radiative transition from the O_{2p} states to the W_{5d} states while the green band is due to the electronic transition from the Ag⁺–V₀ defects 479 level to the VB [63]. Blue/Green emission was also related to distorted [WO₆] octahedral, 480 while red emission to the $[AgO_y]$ (y = 2, 4, 6, and 7) clusters that form complex vacancies, 481 482 inducing more disorder and deeper defects in the forbidden band gap [32,39]. Furthermore, the $[AgO_y]$ (y = 2, 4, 6, and 7) and $[WO_6]$ clusters in pairs have 483 extrinsic defects, which are linked to order-disorder effects in the electronic structure, 484 surface, and interfaces, which create additional energy states above the VB and below the 485 CB, which decrease the band gap [32,67]. However, a specific assignment of surface and 486 487 the clusters that are present on each surface that contributes to PL emission has not been 488 observed. Here, we can observe the influence of particle morphology and surface band gap since the distortions in the lattice favor different distributions of electronic state in 489 490 the surface (Figure 9).

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494 To analyze the difference between the emission spectrum and the contribution of 495 each surface (Figure 9), a comparison among the MAH-8, EI-8, and FI-8 samples with the cube-, rod- and sphere-like morphologies, respectively, is presented in Figure 10. 496 Since each of the surfaces of α -Ag₂WO₄ presents a different band gap energy (E_{gap}^{surf}) 497 498 value, it is expected that each surface has a specific contribution in the PL property. This can be justified by the fact that each surface is quite different in terms of the coordination 499 500 number of atoms, the distance between adjacent atoms, and available electron density [22,30]. 501

Figure 9.

504

505 The theoretical E_{gap} values of surfaces and the rod- and cube-like morphologies of α -Ag₂WO₄ are listed in **Table 1.** According to the results, all surfaces studied show 506 E_{gap}^{surf} values reduced in comparison to the E_{gap} value of the bulk. The density of defects 507 on the surface can justify the existence of intermediary states in the band region of the 508 bulk, which come from the number of Ag-O and W-O bonds that are broken when a 509 slab surface is created. To quantify the defects on the surface, the density of broken bonds 510 was calculated as $D_b = N_b/A$, there N_b is the number of broken bonds and A is the surface 511 512 area. The results corroborate that those surfaces with higher D_b values present the lowest band gap (Table 1) and consequently, intermediary states (Figure 9). 513

Regarding the $E_{gap}^{polyhedron}$ for each experimental morphology, the results reveal 514 that in passing from ideal to the experimental rod-like morphology, the $E_{gap}^{polyhedron}$ value 515 516 increase from 0.99 eV to 1.10 eV due to the appearance of the (101) surface which comes to dominate the Wulff construction (79.4 %) for the rod-like morphology and has a band 517 gap of 1.15 eV. The (010) and (001) surfaces are also exposed in this morphology in a 518 lower proportion (7.3 % and 13.2%, respectively). In the case of the passing from ideal 519 to the experimental cube-like morphology, the $E_{gap}^{polyhedron}$ decrease from 0.99 eV to 0.92 520 521 eV at the time that (001) and (100) surfaces that have the lower band gap values 0.65 eV and 0.48 eV, respectively, increase their contribution (28.6 % each one) to the cube-like 522 morphology that exposed the (010) surface (42.9 %) which have band gap (1.39 eV). 523

It is worth nothing that theoretical values of band gap energy are not the same that those obtained experimentally because the experimental techniques used for such measurements considering the effect bulk/surface of the material and in the calculations have been considered only the exposed surface structures at the morphology.

Figure 10a shows that both CP and EI-8 samples are composed only of rod-like 528 morphologies. The PL spectrum for the CP and EI-8 samples showed two maximum 529 emission centers – around 550 nm and 733 nm – regions. In the case of the MAH-8 530 sample, it is composed of both cube- and rod-like morphologies and although the PL 531 spectrum also present two emission centers, is observed that the maximum emission 532 center around 550 nm is favored while the emission center is 733 nm present intensity 533 lower. Finally, the sample irradiated by femtosecond laser (FI-8) with rod- and sphere-534 like morphologies presented an equilibrium in the maximum emission centers with a 535 slight shift to the red region. Based on these results, we can conclude that (010) and (101) 536 537 surfaces contribute to the emission centers at 550 nm and 733 nm, respectively.

As previously mentioned, (010) surface has the highest band gap, therefore it tends 538 to contribute to the highest energies emission (Figure 10b), in addition, is present in both 539 540 morphologies, see **Table 1**, and is the aim component of the cube-like morphology, thus, the presence of both cube and rod-like morphologies in the sample contributed to the 541 542 emission to 550 nm. In contrast, (101) surface is the main component of the rod-like morphology, and therefore the sample only with rod shows a defined emission at 733 nm. 543 The highest intensity of emission in 550 nm is associated with the presence of localized 544 545 states in the forbidden band gap due to (010) surface, see Figure 9 and Figure 10b, which favored the probability of the transition in this surface. 546

As has been established in the (101) surface are present distorted $[WO_6]_d$ cluster, and the undercoordinate $[AgO_5 \cdot 2V_0^x]$, $[AgO_4 \cdot 3V_0^x]$, and $[WO_5 \cdot V_0^x]$ clusters. The complete distorted $[WO_6]_d$ clusters are considered as a source of electrons due to while the undercoordinate $[AgO_5 \cdot 2V_0^x]$, $[AgO_4 \cdot 3V_0^x]$ clusters, and $[WO_5 \cdot V_0^x]$ clusters supporting charge positive are considered a source of the hole [76,77]. In the case of the (010) surface complete distorted $[WO_6]_d$ and $[AgO_4]_d$ clusters present in the surface are a source of electrons while undercoordinate $[AgO_5 \cdot 2V_0^x]$ clusters are a source of holes, thus, the transfer electron-hole occur from complete distorted clusters to undercoordinate $[AgO_5 \cdot 2V_0^x]$ clusters, the above also explain the contribution of the (010) surface to the PL property [21].

Moreover, the cube- and rod-like morphologies have a stable (001) surface, which has the E_{gap}^{surf} (0.65 eV), see **Table 1** and **Figure 10b**. The (001) surface has a percentage of total area contribution in both morphologies of 28.6% and 13.2%, respectively. The (001) surface has $[WO_5.V_0^x]$, $[AgO_5.2V_0^x]$, $[AgO_4.3V_0^x]$, and $[AgO_4.2V_0^x]$ clusters [21]. The cube-like morphology also has a stable (100) surface, which has the lowest E_{gap}^{surf} (0.48 eV), **Figure 10b**. The (100) surface has $[WO_5.2V_0^x]$, and two $[AgO_5.2V_0^x]$ clusters [21].

In general, the samples with cube-like morphologies showed a maximum center emission in the green region, while those with particles with only rod-like morphology showed two maximum emission centers (in the green and red regions). Differently, the sphere-like morphology presented an equilibrium in the emission centers. The PL behavior demonstrates the influence of many factors, such as the orientation between particles, the variations in the particles size distribution, the morphology of the particles, and surface defects [32].

The XYZ color space created by the International Commission on Illumination (abbreviated as CIE) serves as a standa.rd reference to represent the color emitted by the materials, and this color is expressed by a resulting chromaticity symbolized by the coordinates (x, y) [61]. The chromaticity coordinates x and y under excitation of 355 nm were obtained in the CIE XYZ color space (**Figure S6**), and the detailed information is shown in **Table S4**.

The MAH group revealed variation in their color from orange to red (Figure S6a) 577 578 due to the synthesis time, which disorganized the [AgOy] clusters. The CIE diagram of 579 the EI and FI groups also showed a color variation from orange to red (Figure S6b). Among the samples submitted to femtosecond laser irradiation, the FI-32 was found to 580 be the most resistant to the laser action, remaining practically unchanged when compared 581 to the MAH-32 (Figure S6c and Table S4). The other samples migrated from orange to 582 583 red, following what was observed for samples irradiated by electrons. Therefore, the PL measurements demonstrated that microwave, electron beam, and femtosecond laser 584 irradiations altered the electronic density of the bulk and surface since the behavior of the 585 586 samples changed after irradiation. However, the sample obtained at the longest synthesis 587 time did not show variation in the density of VAg and Vo vacancies when subjected to electron or femtosecond irradiation, which is possibly related to its more stable rod-like 588 589 morphology.

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591 4. CONCLUSION

In summary, in this work, we discussed the influence of morphology, surface band 592 gap, and microstructure on the PL property of α -Ag₂WO₄ samples obtained by the CP 593 method, followed by treatment with MAH at different synthesis times and two different 594 irradiation processes: electron beam and femtosecond laser. The experimental and 595 theoretical study clarifies the important relationship between the DOS of different 596 597 surfaces and the PL property. It was observed that the use of different ways of treating the samples successfully changed the surface characteristics, microstructure, and 598 electronic density of the α-Ag₂WO₄ materials. Electron beam and femtosecond laser 599 irradiations were found to cause expansion of the unit cell, form oxygen vacancies on the 600 surface, change the angle and distance between O-Ag and O-W bonds, and modify the 601

particle morphology to rod-, cube- and sphere-like. The theoretical calculations showed 602 603 how changes in the microstructure, morphology, and surface band gap values can alter the distribution of electron density, and consequently the emission centers. Cube-like 604 morphology shows a high contribution of the (010) surface, while rod-like morphologies 605 showed the contribution of (010) surface and (101) surface. Moreover, the sample 606 607 submitted to a longer synthesis time of 32 min in the microwave system presented the 608 most stable rod-like morphology and high resistance to the disorder of the networkmodifying $[AgO_v]$ clusters. Therefore, the more ordered the network-forming $[WO_6]$ 609 610 clusters, the more difficult it will be to introduce defects. For this reason, no significant 611 changes were observed in the PL property. These results enrich the literature regarding the effect of different techniques associated with microstructure, morphology, and band 612 gap energies on the PL properties of semiconductor materials. 613

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646 NOTES

647 The authors declare no competing financial interest.

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954 FIGURE CAPTIONS

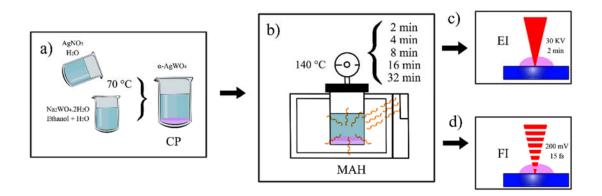
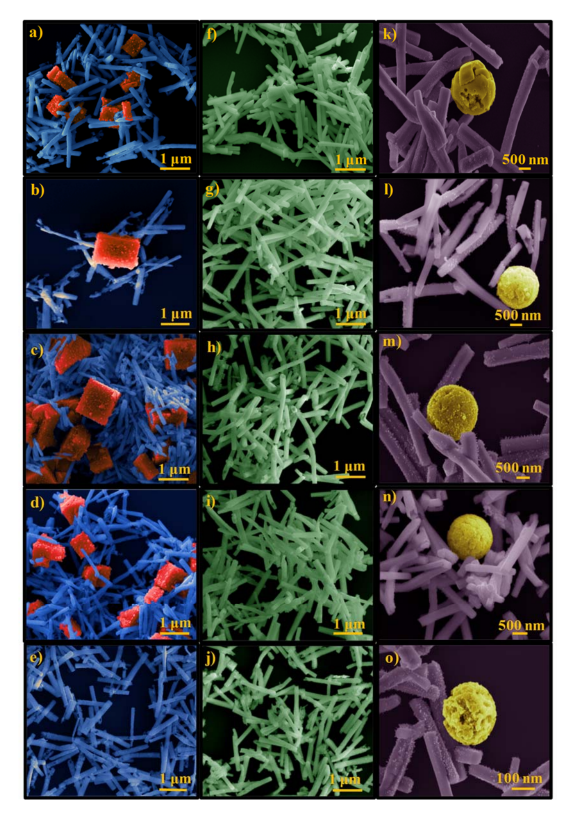


Figure 2. Scheme with steps of a) Synthesis by CP method, b) Treatment by MAH
method, c) Electron beam irradiation, and d) Femtosecond laser irradiation; to obtain αAg₂WO₄ samples.

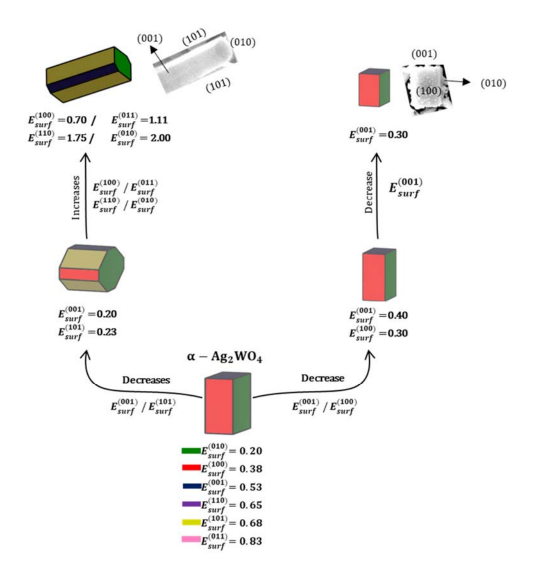




963 Figure 2. FE-SEM images of α -Ag₂WO₄ particles of MAH, EI, and FI. (a-e) 964 heterogeneous morphology obtained after treatment with MAH at different treatment 965 times (2, 4, 8, 16, and 32 min) with the cubes highlighted in orange color and the rods-966 like in blue color; (f-j) rods-like with homogeneous morphology after EI indicated by

967 green color; (k-o) heterogeneous morphology of rods in violet color and spheres in yellow

968 color after FI.



972 Figure 3. Wulff construction for α -Ag₂WO₄. For comparison, experimental FE-SEM

973 images are shown. E_{surf} values are given in Jm^{-2} .

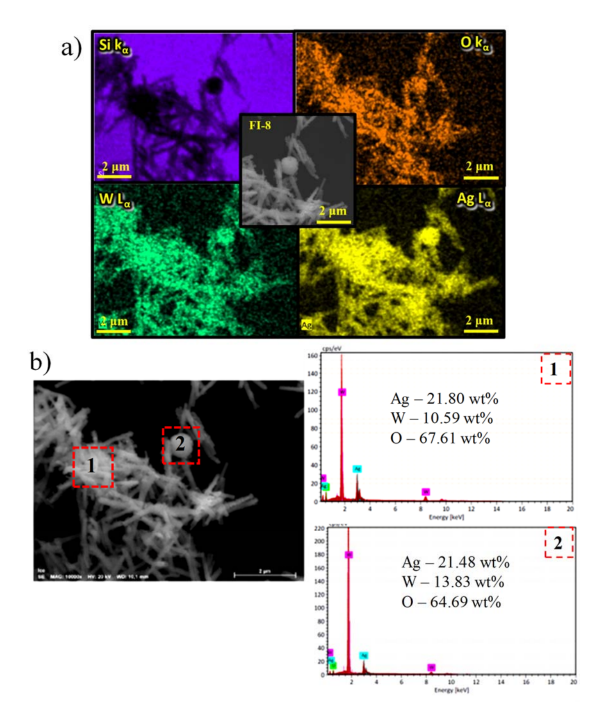
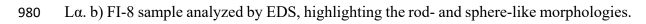
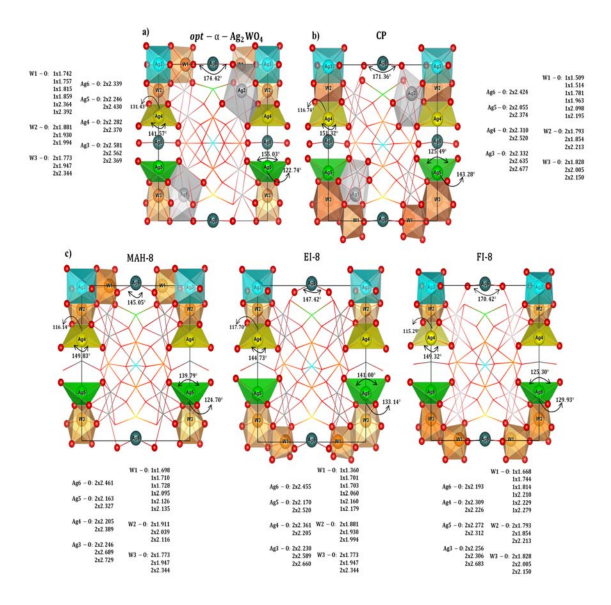


Figure 4. a) EDS elemental mapping of the FI-8 sample of Si K α , O K α , W L α , and Ag



- 981 c) Rod-like region 1 and d) Sphere-like region 2.



986 Figure 5. Geometry of a) $opt - \alpha$ -Ag₂WO₄, b) CP, and c) irradiated (MAH-8, EI-8, and

- 987 FI-8) structures.

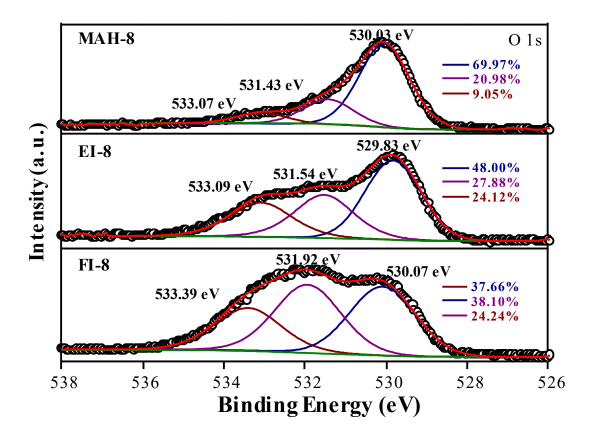
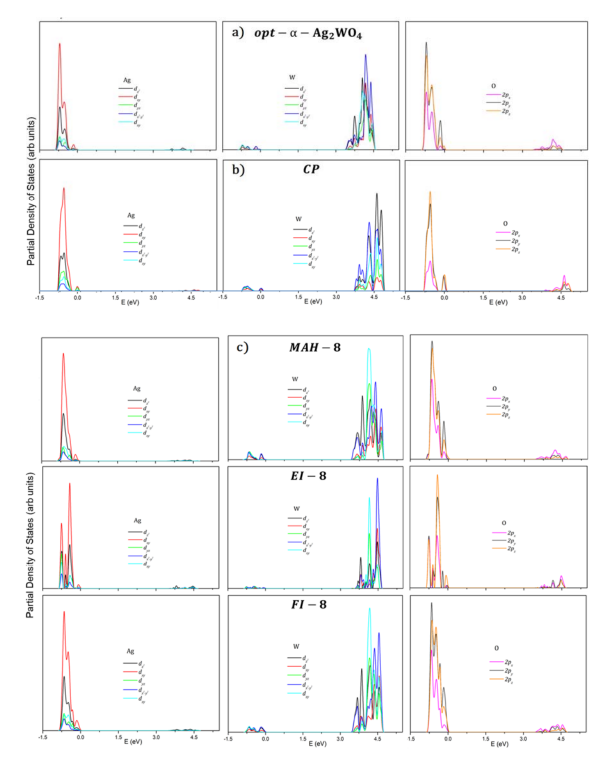
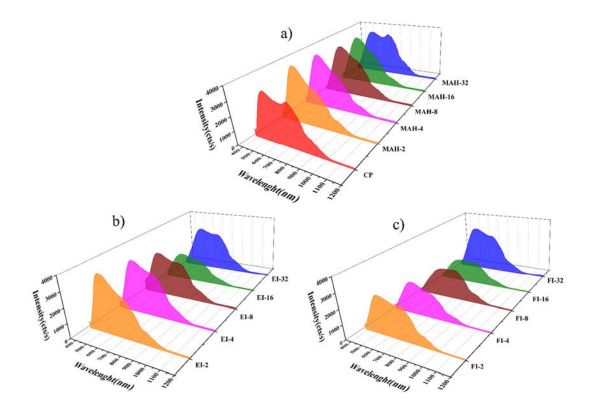


Figure 6. High-resolution XPS spectra of O *ls* for the MAH-8, EI-8, and FI-8 samples.



997

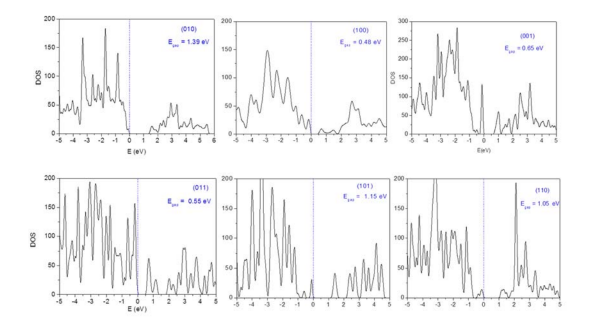
Figure 7. Partial density of states projected on the 4*d*, 5*d* and 2*p* orbitals of Ag, W, and O atoms, respectively, for a) $opt-\alpha$ -Ag₂WO₄, b) CP, and c) irradiated (MAH-8, EI-8, and FI-8) samples.





1004 Figure 8. PL spectra of the samples obtained by a) the CP method, treated with MAH

- 1005 and irradiated by b) EI and c) FI.





1011 Figure 9. Total density of states for the (010), (100), (001), (011), (101), and (110)

- 1012 surfaces of α -Ag₂WO₄.

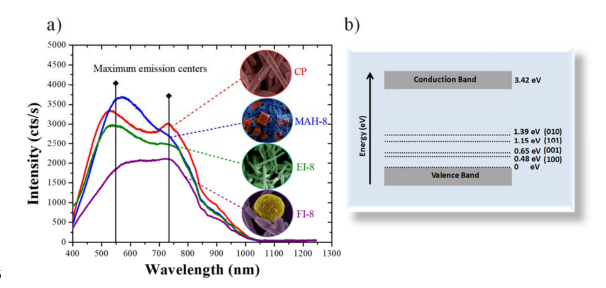


Figure 10. (a) PL spectra and maximum emission centers of the CP, MAH-8, EI-8, and
FI-8 samples. (b) Comparative diagram of the band gap value of the optimized structure
(3.42 eV) and band gap values for (100), (010), (001), and (101) surfaces.