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A Systematic Study of the Effect of Silver on the Chelation of Formic Acid to a Titanium Precursor and the Resulting Effect on the Anatase to Rutile Transformation of TiO₂

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Anatase to rutile transition in an unmodified synthetic titania usually occurs at a temperature range of 600-700°C. Various methods such as addition of metallic and nonmetallic dopants and modifying the precursor have previously been reported to influence the anatase to rutile transition temperature. In the current study, the effect of addition of increasing amounts of silver to the extent of chelation of a formate group to a titanium precursor and the resulting effects on the transformation of anatase to rutile has been studied. The addition of silver (0, 1, 3, and 5 mol %) on the anatase to rutile transformation temperature has been systematically followed by Fourier transform infrared (FTIR), Raman, X-ray diffraction (XRD), differential scanning calorimetry, and X-ray photoelectron spectroscopy (XPS) studies. From the FTIR and Raman spectroscopy studies it was observed that the incorporation of silver caused a reduction in the intensity of the COO⁻ stretches indicating that the titania formate bridging complex is becoming weaker in the presence of silver. XRD studies indicated an early rutile formation for the silver-doped samples. XRD of the samples calcined at 700 $^{\circ}$ C showed that 5 mol % Ag TiO₂ contained both anatase (46%) and rutile (54%), whereas the undoped sample primarily consists of anatase (95%). At 800 °C all silver doped samples converted to 100% rutile and the undoped TiO₂ consisted of both anatase (55%) and rutile (45%). XPS analysis showed that Ag^0 and Ag_2O has been formed on the surface of the titania formate complex without calcination (>100 °C) indicating that photo-oxidation has occurred. FTIR, Raman, and XPS studies confirmed that the presence of silver in the xerogel before calcination may be responsible for the reduction of the titanium formate bridge. It was concluded that the presence of silver (Ag₂O and Ag⁰) hindered bridging ligands, which resulted in a weakened titanium gel network. This structurally weakened gel network could easily be collapsed during calcination, and it favors an early rutile formation.

30 Introduction

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31 Titanium dioxide semiconductor photocatalysis has attracted 32 the attention of several researchers in the past decade due to its environmental applications such as air purification and water 33 remediation.¹⁻¹⁰ Anatase, rutile, and brookite are the three 34 35 polymorphs of TiO₂ and differ only in arrangement of their [TiO₆]²⁻ octahedra; anatase (tetragonal) consists of octahedrals 36 sharing vertices; rutile (tetragonal) is connected by edges, and 37 in brookite (orthorhombic) both edges and vertices are 38 connected.^{11–13} Rutile is found freely in nature, but all three 30 can be synthetically prepared. Rutile is the thermodynamically 40 stable phase, while anatase and brookite are both metastable, 41 transferring to rutile under heat treatment at temperatures 42 typically ranging between 600-700 °C.11 Anatase is widely 43 regarded as the most photocatalytically active of the three 44 crystalline structures.^{13–15} Titanium dioxide is the most widely 45 investigated photocatalyst because of its ease of preparation, 46 availability, strong oxidizing power, nontoxicity, and long-term 47

stability.^{7,16} However, because of its large band gap (3.2 eV for 48 anatase), TiO₂ can only be activated upon irradiation with a photon of light <390 nm, limiting its use under solar irradiation.^{16–18} Because of this, researchers have been focusing their attention on ways to improve the photocatalytic efficiency of TiO₂ under irradiation with visible light (>400 nm). 53

Asahi et al. have reported nitrogen doped TiO₂ promoting 54 photocatalytic activity up to $\lambda = 520$ nm claiming that the 55 presence of nitrogen narrows the band gap of TiO₂ thus making 56 it capable of performing visible light driven photocatalysis.¹⁹ 57 However, Ihara et al. suggested that it is the oxygen vacancies 58 that contributed to the visible light activity, and the doped 59 nitrogen only enhanced the stabilization of these oxygen 60 vacancies.²⁰ They also confirmed this role of oxygen vacancies 61 in plasma-treated TiO2 photocatalysts.²⁰ In addition the structural 62 oxygen vacancies causing visible light photocatalytic activity 63 was also reported by Martyanov et al.²¹ Further studies with 64 nonmetal dopants, S², C², I², Br, and Cl¹⁸ also show red shifts 65 in band gap edge of TiO_2 . 66

Transition metal doping has also given promising results for 67 visible light activated TiO₂ by extending the absorption spectra into the visible region. Much research has focused on the transition metal ion Fe³⁺²⁵⁻²⁹ whereby its incorporation into the results in the formation of new energy levels 71

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between the valence band and the conduction band.³⁰ Deposition of noble metals Ag, Au, Pt, and Pd on the surface of TiO_2 enhance the photocatalytic efficiency by acting as an electron trap, promoting interfacial charge transfer and therefore delaying recombination of the electron—hole pair.^{27,31–34}

Many researchers have focused on modifying TiO₂ with Ag. 77 For example, Chao et al. reported the effect of Ag doping on 78 the phase transformation and grain growth of sol-gel TiO₂ 79 powder.³⁵ Kuo et al. showed through X-ray diffraction (XRD) 80 and X-ray photoelectron spectroscopy (XPS) that silver on TiO_2 81 surface coatings was easily oxidized into silver oxide (Ag₂O) 82 and that the addition of silver causes a reduction in photolu-83 minescence (PL) intensity as found by PL spectroscopy.³⁶ This 84 group previously showed enhanced visible light photocatalysis 85 with Ag modified TiO2.37 Choi et al. controlled the ratio of 86 87 anatase and rutile phases through the addition of surfactants.³⁸ 88 The effect of precursor chelation on anatase to rutile transition has also been reported previously. Acetic acid,³⁹ formic acid,⁴⁰ 89 urea,⁴¹ sulfuric acid,⁴² and ammonium sulfate⁴³ have previously 90 been employed to study the effect of a chelating agent on the 91 anatase to rutile transformation in the TiO₂ photocatalyst. 92 However, there are no systematic studies on the effect of silver 97 doping on chelation and anatase to rutile transformation. 94

95 The current paper reports a systematic study on how the addition of increasing amounts of silver affects the extent of 96 97 chelation of a formate group to a titanium precursor and how the resulting reduction in formate chelation causes early 98 transformation of anatase to rutile. The effect of the addition 99 of silver on structural changes is investigated by characterizing 100 the sample in its amorphous state (i.e., before it is calcined) 101 with XPS, IR, and Raman studies. XRD was used to determine 102 the crystalline phase of the calcined samples, and differential 103 104 scanning calorimetry (DSC) was employed to examine the 105 thermal events of the sample through heat treatment. The effect 106 of silver on the electronic transitions of crystalline TiO₂ is shown through UV-vis and PL spectroscopy. To the best of the 107 authors' knowledge this is the first paper to report the effect of 108 silver on a titanium formate complex before calcination and 109 the subsequent effects of the presence of silver upon calcination. 110

111 Experimental Section

Titanium tetraisopropoxide (TTIP)(97%), formic acid (98%), 112 and silver nitrate (99%) were purchased from Aldrich and used 113 without further purification. Deionized water was used in all 114 experimental preparations. The samples were prepared by a 115 modified sol-gel route.⁴⁰ Titanium tetraisopropoxide (36 mL) 116 was added to formic acid (19 mL) under stirring. Water (9 mL) 117 was added, and a thick paste was formed. The TTIP, formic 118 119 acid, and water were used in 1:4:4 molar ratios. The mixture 120 was stirred for 2 h and filtered, and the filtrate was dried in an oven at 100 °C for 12 h. To prepare silver doped titania, the 121 above procedure was repeated, including silver nitrate (1, 3, 122 and 5 mol %) in the water before it was added to the TTIP/ 123 formic acid mixture. The dried powders were calcined at 300, 124 500, 700, and 900 °C for 2 h at a ramp rate of 3 °C/min. A 125 Siemens D 500 X-ray diffractometer, with a diffraction angle 126 range $2\Theta = 20-80^{\circ}$ using Cu K α radiation, was used to collect 127 XRD diffractograms. The mass fraction of rutile (X_R) in the 128 calcined samples was calculated using the Spurr equation⁴⁴ (eq 129 1) 130

$$X_{\rm R} = \frac{1}{1 + 0.8(I_{\rm A}/I_{\rm R})} \tag{1}$$



Figure 1. XRD of 0 and 5% Ag TiO₂ powders calcined at 700 °C.

where I_A is the main intensity of anatase (101) peak and I_R is 131 the main intensity of rutile (110) peak. 132

The crystallite size (*T*) was estimated using the Scherrer 133 equation³⁷ (eq 2) 134

$$T = \frac{0.9\lambda}{\beta\cos\theta} \tag{2}$$

where T is the crystalline size, λ is the X-ray wavelength, θ is the Bragg angle, and β is the line broadening.

A Perkin-Elmer Lambda 900 UV-vis absorption spectro-137 photometer was used to record absorption and diffuse reflectance 138 spectra; samples were mixed in KBr (1:20 sample KBr) and 139 pressed into a tablet; a KBr tablet made under the same 140 conditions was used as a reference. IR spectra were obtained 141 using a Perkin-Elmer GX FTIR spectrometer and recorded as a 142 KBr disk (1:10 sample/KBr); Raman spectra were recorded on 143 an ISA Labram, employing an argon laser (514.5 nm) as an 144 excitation source. PL was recorded on a Perkin-Elmer LS55 145 luminescence spectrometer using an excitation wavelength of 146 320 nm. Approximately 5 mg of sample was placed into an 147 aluminum sample pan for DSC using an empty aluminum pan 148 as a reference. All DSC were recorded on a Shimadzu DSC-60 149 between 25 and 600 °C at a ramp rate of 20 °C/min. 150

Results and Discussion

XRD. XRD was carried out on the calcined samples in order 152 to determine the crystalline phase of the samples. All samples 153 calcined at 300 °C were amorphous. Crystalline anatase TiO₂ 154 was present for all samples calcined at 500 and 600 °C. 155 However, at 700 °C the diffractogram (Figure 1) showed that 5 156 mol % Ag TiO₂ contained both anatase (46%) and rutile (54%), 157 but TiO₂ without silver consisted mainly of anatase (95%). This 158 indicated that the presence of increased amounts of silver 159 promotes the anatase to rutile transformation. 160

Promotion of phase transformation by the addition of silver 161 is believed to be caused by the following factors.³⁵ Decreasing 162 anatase grain size (Figure 2) results in an increase in the total 163 F2 boundary energy for the TiO₂ powder. The driving force for 164 rutile grain growth is therefore increased, which promotes 165 anatase to rutile phase transformation.⁴⁵ As the transformation 166 of anatase to rutile is a mechanism of nucleation and growth,^{46,47} 167 an increased amount of nucleation sites would favor rutile 168 formation. Phase transformation is also governed by such effects 169 as defect concentration⁴⁸ and grain boundary concentration,⁴⁹ 170

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Figure 2. Variation in the nanocrystallite size (from XRD) as a function of increase in concentration of silver, at different calcination temperatures.



Figure 3. XRD of 0 and 5% Ag TiO₂ powders calcined at 800 °C.

the presence of which can be expected to be increased with
greater surface areas. Rutile nucleation is thus enhanced as the
presence of defect sites is increased. Therefore, an increase in
the number of defect sites promotes anatase to rutile transformation at lower temperatures.³⁵

176 At 800 °C, the undoped TiO₂ sample consisted of both anatase 177 (55%) and rutile (45%), whereas the silver-doped samples were 178 all 100% rutile (Figure 3). A higher Ag concentration promotes 179 earlier phase transformation. Figures 1 and 3 show that the 180 presence of silver promotes the formation of rutile.³⁵

The radius of Ag^+ ion (126 pm) is much larger than that of 181 the Ti⁴⁺ ion (68 pm) and so Ag⁺ ions cannot enter the lattice 182 of anatase TiO2.^{35,37} However, migration of the Ag⁺ ions from 183 bulk anatase grains to the anatase grain surface can occur during 184 calcination.35,37 With Ag migrating to the TiO2 surface, surface 185 defects in the anatase grains will increase. This results in a 186 187 greater number of nucleation sites for the formation of rutile, again, promoting phase transformation. Oxygen vacancies may 188 also influence the anatase to rutile transformation.^{35,50,51} Previous 189 reports^{50,51} indicate that the concentration of oxygen vacancies 190 at the surface of anatase increases with Ag doping. This favors 191 the ionic rearrangement necessary for the structure changes 192 associated with rutile phase formation.35 193

An alternate effect that the silver may have on early rutile formation is that Ag cations may be easily reduced. The reduced



Figure 4. IR spectra of TiO_2 powders with different mol % silver content after aging at 100 °C.

 Ag^+ ions can spread on the anatase surface to Ag^0 by heat. Photo 196 reduction of the Ag⁺ ions may also occur as the samples were 197 never protected from light irradiation.^{52,53} Oxygen vacancies will 198 occur for charge compensation caused by Ag⁺ reduction, and 199 again these oxygen vacancies are favored for the formation of 200rutile.35,50,51 To further investigate the role silver plays in altering 201 the transformation temperature of TiO₂ in this study, it is 202 necessary to understand the structure of the oligomer before 203 calcination. To gain insight on this, DSC, IR, Raman, and XPS 204 were carried out on the powders before calcination. 205

IR Spectroscopy. The influence of silver on the structure of the TiO powders was investigated before calcination using IR and Raman spectroscopy. Figure 4 shows IR spectra of the doped and undoped powders. Symmetric and asymmetric COO⁻ stretches at \sim 1360 and 1540 cm⁻¹, respectively, indicated that the formic acid forms a bidentate bridge with the titanium precursor.^{40,54-56} 212

Figure 4 also shows that silver clearly reduces the intensity 213 of the COO⁻ stretches as with each increase in Ag concentration 214 there is a reduction in intensity of both the asymmetric and 215 symmetric COO⁻ peaks. This result provides evidence for an 216 alternative mechanism to those stated above for early anatase 217 to rutile transformation in the presence of silver. In fact, silver 218 may influence the anatase to rutile transformation through 219 interactions with the titanium precursor in the early stages of 220 the sol-gel synthesis. These interactions may ultimately alter 221 the condensation pathway, resulting in a weakened TiO_2 222 oligomer network. 223

To compare the effects of aging at 100 °C, IR spectra of samples 0 and 5 mol % silver TiO₂ were recorded before and after aging (Figures 5 and 6).

After both samples are aged at 100 °C for 12 h there is a significant reduction in the bridging formate COO⁻ stretches 228 of the 5% silver doped sample, whereas the TiO₂ sample without 229 silver does not show a significant reduction in the carboxylate 230 stretches. Since the boiling point of formic acid is 101 °C, it is 231 possible that the formic acid is displaced by the presence of 232 silver and is then evaporated upon aging. 233

Raman Spectroscopy. Raman spectroscopy analysis was234carried out on the powders before they were calcined to support235IR spectroscopy results.236

The Raman spectra of the doped and undoped TiO_2 samples, 237 (Figure 7) supports the IR results. As the silver content is 238 increased, the asymmetric and symmetric COO⁻ stretches (1570 239 and 1390 cm⁻¹) decrease accordingly. The addition of silver 240

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225 226 **F5**

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Figure 5. IR spectra of TiO_2 powders with different mol % silver content before aging.



Figure 6. IR spectra of TiO₂ powders with different mol % silver content after aging at 100 °C for 12 h.



Figure 7. Raman spectra of TiO_2 powders with different mol % silver content before calcination.

also causes significant changes in the TiO region of the spectra (0-1000 cm⁻¹). The presence of peaks at 160, 420, 515, and 620 in the Raman of the silver doped TiO₂ are indicative of the four peak pattern that would be expected for anatase.⁵⁷⁻⁵⁹ However, without the presence of silver, this four-peak pattern



Figure 8. DSC of 0 and 5% Ag before aging.



Figure 9. DSC of 0 and 5% Ag after aging at 100 °C.

is not as distinctive, again providing further evidence that silver effects the titanium formate complex before crystallization has occurred. 248

It is agreed that the use of a chelating agent gives stability to 249 the hydrolysis and condensation reactions associated with the 250 sol-gel process in the production of TiO₂ from titanium 251 alkoxides.54,60-64 IR and Raman spectroscopy have shown that 252 formic acid forms a bridging ligand with titania.55,65-68 Previous 253 reports have shown that similar chelating agents remain bound 254 to the central titanium atom, while the isopropoxy (OR) groups 255 are preferentially hydrolyzed. The bridging ligands remain 256 throughout much of the condensation process, ^{54,56} altering the 257 condensation pathway and promoting the formation of linear 258 polymers composed of edge sharing octahedral.^{39,56} The system 259 may be destabilized through the addition of water, which leads 260 to a structurally weak network that can easily collapse upon 261 calcination to form rutile.69,70 262

From the spectroscopy results it can be seen that the addition of silver causes a reduction in the intensity of the COO⁻ stretches indicating that the titania formate bridging complex is becoming weaker in the presence of silver. This may lead to a structurally weak oligomer that upon calcination easily forms rutile at lower temperatures.

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DSC. DSC was carried out to investigate the thermal events associated with the doped and undoped TiO_2 samples before and after aging.

The DSC curve of the undoped TiO_2 powder analyzed before 272 aging (Figure 8) is almost identical to the same sample after 273 aging (Figure 9). It reveals an endothermic peak at ~100 °C 274 attributed to the elimination of unbound water and formic acid 275 from the surface of the TiO_2 powder. The same peak for the 276 silver doped sample reveals as expected, a smaller enthalpy of 277

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Figure 10. XPS spectra of C 1s (a) and O 1s (b) of TiO_2 with and without silver before calcination.



Figure 11. XPS spectra of Ti 2p (a) and Ag 3d (b) of 3% Ag TiO₂ before calcination.

-420 J/g compared with -636 J/g for the undoped sample,
indicating that formic acid is easier to remove in the presence
of silver. This is in agreement with the previously proposed
mechanism from the FTIR and Raman spectroscopy results, that
silver inhibits formic acid and it is thus removed upon aging at
100 °C therefore, allowing the oligomer structure to readily
collapse and form rutile upon calcination.

285 In Figures 8 and 9 an endothermic peak at \sim 250 °C is due to the removal of isopropanol formed through the condensation 286 287 step. The removal of isopropanol indicates that the condensation step is complete. Comparing the DSC curves of the samples 288 before (Figure 8) and after (Figure 9) aging shows that the 289 endothermic isopropanol peak is present in both the 0 and 5 290 mol % silver sample before aging. But after aging the isopro-291 panol peak at 250 °C is only present with 0% Ag TiO₂. This 292 provides further explanation for the formation of rutile at lower 293 temperatures, since when the condensation step is near comple-294 tion, the crystallization temperature is lowered and then so too 295 is the anatase to rutile transformation temperature. 296

The appearance of the first exothermic peak in all DSC curves 297 is the transition from amorphous TiO₂ to crystalline anatase, 298 and as expected this peak occurs earlier with silver doped TiO₂ 299 than with TiO₂ only. Figure 8 shows that crystallization occurs 300 301 at 320 °C for 5% Ag TiO₂ but does not occur until 350 °C for the undoped sample. This can also be seen in Figure 9 where 302 crystallization of 5% Ag TiO₂ takes place at 320 °C but for the 303 undoped sample crystallization does not occur until 385 °C. 304

305 XPS. To determine more information on exactly how the
 306 silver is interacting with the titanium formate complex, XPS
 307 was carried out on the silver doped and undoped samples before

calcination to establish the titanium structure, the chemical state 308 of the silver particles, and also for further evidence of the 309 reduction in the carboxylate species. XPS spectra were recorded 310 of the samples precalcination (as with IR and Raman) and XPS 311 was also carried out on the crystalline titania after calcination. 312 The spectra of C1s and O1s of TiO₂ without silver show the 313 presence of the carboxylate group at 289.1⁷¹ and 532.3 eV,^{71,72} 314 respectively (parts a and b of Figure 10). As the silver content 315 is increased the intensity of these peaks decreases in a similar 316 manner as those of the IR and Raman results, again indicating 317 that with increased silver content, the formation of a titania 318 formate bridging complex becomes increasingly difficult to 319 form. 320

Figure 11 shows the XPS narrow scans for Ti 2p and Ag 3d 321 peaks. The XPS spectrum of Ti 2p is unchanged with increasing 322 amounts of silver, the Ag 3d spectrum of 1% silver TiO2 gives 323 a weaker signal than the 3% shown in figure 11b and the Ag 324 3d scan of 5% silver TiO₂ results in a spectrum similar to 3% 325 silver TiO₂. The fact that increasing amounts of silver does not 326 affect the Ti 2p spectra indicates that central titanium atom has 327 not been reduced. In Figure 11a, a Ti 2p peak at 459.1 eV is 328 representative of Ti in its tetravalent state⁷³ in an octahedral 329 environment.⁷⁴ The absence of a Ti³⁺ peak at 457.4 eV leads 330 to the following observations. 331

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(1) Ti has not been reduced to Ti^{3+} which may indicate an 332 absence of surface oxygen vacancies.⁷⁵ However, if TiO_n (n < 333 2) is formed it may not be present in detectable amounts.⁷⁶ 334

(2) Ag_2O or Ag^0 incorporation into the TiO₂ lattice may give 335 rise to a signal at 457.4 eV representative of Ti³⁺.³⁶ From the 336

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Figure 12. O 1s XPS spectra of TiO₂ without silver calcined at 900 °C (a) and with 3% silver calcined at 700 °C (b).



Figure 13. Ti 2p XPS spectra of TiO₂ without silver calcined at 900 °C (a) and with 3% silver calcined at 700 °C (b).

337 XPS there was no evidence of Ag TiO_2 bond formation, which 338 can be expected due to the differences in atomic radius.

The Ag 3d scan (Figure 11b) shows two large peaks. A 339 340 Gaussian fit of the main peak (\sim 368 eV) showed that it was made up of two signals at 367.9 and 368.2 eV, representing 341 the chemical bonding states of Ag₂O and Ag⁰, respectively.³⁶ 342 Therefore, it has been shown that Ag⁰ and Ag₂O have formed 343 344 on the surface of the titania formate complex, before heat treatment above 100 °C. Spectroscopic (FTIR, Raman, and XPS) 345 studies of the carbonyl species have shown a reduction in the 346 titanium formate bridging complex with increased amounts of 347 silver. The Ag 3d scan of a sample (before calcination) 348 containing silver (Figure 11b) shows the presence of Ag₂O 349 $(Ag^{2+}, 367.8 \text{ eV})$ and $Ag^0 (368.2 \text{ eV})$.³⁶ Therefore, the presence 350 of Ag₂O and Ag⁰ in the powders before calcination may be 351 responsible for the reduction of the titanium formate bridge as 352 shown by IR, Raman, and XPS. The presence of Ag_2O and Ag^0 353 may then restrict the formation of a titanium formate bridging 354 complex which leads to an altered condensation pathway and 355 therefore low temperature formation of rutile. 356

Figure 12 shows the narrow scan XPS spectra for O 1s of F12 357 358 TiO₂ only calcined at 900 °C and 3% Ag TiO₂ calcined at 700 359 °C. Gaussian fits of both spectra give rise to two peaks at 530.1 and 531.1 eV for crystal lattice oxygen and hydroxy oxygen, 360 respectively.^{74,77} The silver doped sample contains a greater 361 amount of hydroxy oxygen. Chemisorbed surface hydroxyl 362 groups can enhance photocatalysis by trapping photoinduced 363 holes resulting in an increase in the formation of the highly 364 oxidizing OH[•] radicals.^{77,78} It is well reported that silver retards 365 the recombination of photogenerated electron hole pairs;^{36,37,79} 366 this may not only be due to the attraction of excited electrons 367

to silver but also due to the presence of extra hydroxyl species 368 to delay recombination through hole trapping. 369

There is an apparent difference between the Ti 2p spectra 370 shown in Figure 13. Unmodified TiO₂ gives a signal at 458.9 371 eV in the Ti 2p narrow scan. However, 3% silver TiO₂ results 372 in the presence of an additional signal at 459.2 eV. The undoped 373 sample after calcination at 900 °C gives a symmetrical peak at 374 458.9 eV (Figure 13a) typical of tetravalent Ti-O bond.⁷³ 375 However, the Ti 2p narrow scan of the 3% Ag TiO₂ sample 376 calcined at 700 °C does not give a symmetrical peak. A Gaussian 377 fit of the spectrum gives two signals at 458.9 and 459.2 eV 378 both representing Ti-O,⁷³ but the shift in the binding energy 379 to 459.2 eV indicates that there may be an interaction between 380 TiO₂ and silver.⁸⁰ 381

Through XPS, IR, and Raman it was shown that the addition 382 of silver restricts the formation of a titanium formate bridging 383 complex. XPS also shows that silver exists as both Ag⁰ and 384 Ag₂O. Finally, XPS has shown that the presence of silver results 385 in a peak being present at 459.2 eV in the Ti 2p scan (Figure 386 13b), which is indicative of an interaction between TiO_2 and 387 silver.⁸⁰ The combination of these results leads to the proposal 388 of Scheme 1. In Scheme 1 it is proposed that Ag⁰ and Ag₂O 389 block the formation of the titanium formate bridge. The resulting 390 titanium complex can then collapse readily upon calcination to 391 form rutile. 392

Gaussian fits of the narrow scan Ti 2p XPS spectra (Supporting Information) of all samples before calcination reveals two signals. The intensity of the signals accordingly varies with the increasing presence of silver, further indicating that $Ag^{0/3}$ and AgO is interacting with titanium,⁸¹ thus facilitating the removal of formate species and allowing for the collapse of the Ti–O 398

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399 gel framework upon calcination to form rutile at lower 400 temperatures.

401 **Diffuse Reflectance Spectroscopy.** To estimate the band gap 402 distance, UV–vis spectroscopy was employed. The results show 403 that silver does improve visible light absorbance of TiO_2 due 404 to silver plasmon absorption; moreover, a blue shift was 405 observed for the band gap separation of the TiO_2 materials upon 406 increased silver addition.

- F14 407 Figure 14 shows the diffuse reflectance spectra for 0, 1, 3,
 408 and 5 mol % Ag TiO₂. It can be seen that unmodified TiO₂ has
 409 a smaller band gap than the silver modified TiO₂. The blue shift
 410 of the silver modified materials can be attributed to the presence
 411 of additional silver.⁸²
- Figure 15 shows a diffuse reflectance spectrum of Ag F15 412 413 nanoparticles where a strong absorbance is observed at \sim 335 414 nm. The band gap of the TiO₂ nanomaterials can therefore not be accurately determined because of the strong silver absorp-415 tion.⁸³ It is also clear that the presence of increased amounts of 416 silver (3 and 5 mol %) facilitates visible light absorbance. This 417 can also be seen from Figure 15 through strong visible light 418 absorption of the silver nanoparticles. The proposed mechanism 419 for the visible light absorbance of Ag TiO₂ is shown in Figure 420 16.80 **F16** 421
- F17 422 It can also be seen in both Figures 14 and 17 that 3 mol %423 silver TiO₂ has greater absorption than 5 mol % Ag in the visible 424 region. This is because increased levels of silver act as a physical 425 block against TiO₂ light absorption. This causes an increase in 426 the diffuse light reflectance of the material.³⁷
- **F18** 427 In Figure 18 it is observed that the band gap of the modified 428 TiO₂ is reduced, as seen by the red shift in the spectra of the 3 429 and 5 mol % Ag TiO₂ in comparison with Figures 14 and 17. 430 This is attributed to the formation of rutile. Rutile has a smaller 431 band gap than anatase;¹³ XRD results confirm that silver doped 432 samples calcined at 800 °C have rutile structure (Figure 3). By



Figure 14. Diffuse reflectance spectra of 0, 1, 3, and 5% Ag TiO₂ calcined at 500 $^{\circ}$ C.



Figure 15. Diffuse reflectance spectrum of silver nanoparticles.



Figure 16. Mechanism for light absorption of silver.



Figure 17. Diffuse reflectance spectra of 0, 1, 3, and 5% Ag TiO₂ calcined at 700 $^{\circ}$ C.

comparison of Figure 18 with Figures 14 and 17, silver does433not have the same influence in causing the blue shift. The434mechanism of anatase to rutile transformation is one of435nucleation and growth;46,47therefore, rutile particles are sig-nificantly larger than those of anatase. The larger particles may437result in a significant reduction in the influence of silver on the438band gap of the materials.439

Conclusions

A systematic study of the effect of silver on the anatase to rutile transformation temperature of TiO_2 has been carried out. By use of XRD, FTIR, Raman, DSC, and XPS it was proposed that the addition of silver blocks the formation of a titanium -

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Figure 18. Diffuse reflectance spectra of 0, 1, 3, and 5% Ag TiO₂ calcined at 800 $^{\circ}$ C.

carboxylate bridging ligand. This was clearly shown through 445 446 the carboxylate stretches in FTIR, Raman and XPS. Without the formation of this bridging ligand the condensation pathway 447 is altered, and the resulting titania polymer network is weakened. 448 When calcined, this weakened structure can readily transform 449 from anatase to rutile due to a greater atomic mobility. The 450 sample with no silver present maintained anatase at greater 451 temperatures than those that contained silver. This was due to 452 the formation of a strong carboxylate bridge that promoted a 453 more organized structure throughout the condensation process. 454 455 The more ordered oligomer network of the sample without silver consisted of anatase at greater temperatures than those where 456 silver was present. This was clearly seen from the XRD 457 diffractograms. Previous reports have indicated that oxygen 458 459 vacancies contribute to the early formation of rutile, but through 460 XPS and PL, it was concluded that the presence of additional silver did not form extra oxygen vacancies. XPS also showed 461 that Ag₂O and Ag⁰ was present in the samples before high 462 temperature calcination. It was also showed through XPS that 463 464 there may be an interaction between Ag and Ti, which agrees with the proposed mechanism outlined in Scheme 1. 465

PL studies showed that the addition of silver reduced
recombination of electron, hole pairs, and PL spectra did not
provide any evidence for the presence of additional oxygen
vacancies with increased amounts of silver. Diffuse reflectance
showed greater visible light absorbance through silver plasmon
resonance.

An alternative mechanism has been proposed on how silver effects the anatase to rutile transition of TiO₂. In the proposed mechanism, silver blocks the formation of a bridging ligand with the titanium alkoxide precursor. This is clearly shown in FTIR, Raman, and XPS spectra, and the resulting lower formation of rutile is clear from XRD and DSC.

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484 Supporting Information Available: Figures depicting Ti
485 2p XPS spectrum of 0, 1, 3, and 5% Ag before calcination and
486 5% silver loading on TiO₂ calcined at 700 and 800 °C. This

material is available free of charge via the Internet at http:// 487 pubs.acs.org. 488

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