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2009-09-03

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Nolan, N., Pillai, S., Seery, M. (2009): Spectroscopic Investigation of the Anatase-to-Rutile Transformation of Sol-Gel Synthesised TiO2 Photocatalysts. *Journal of Physical Chemistry C*, 113, 2009, pp. 16151-16157. doi:10.1021/jp904358g

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Funder: Environmental Protection Agency, Ireland

J. Phys. Chem. C XXXX, xxx, 000

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Spectroscopic Investigation of the Anatase-to-Rutile Transformation of Sol-Gel-Synthesized TiO₂ Photocatalysts

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Received: May 11, 2009; Revised Manuscript Received: July 26, 2009

Among the three major phases in titania, anatase is reported to be a better photocatalytically active phase. Anatase to rutile transformations, under normal conditions, usually occurs at a temperatue range of 600–700 °C. Various chemical additives have previously been employed to extend the anatase transition to higher temperatures. The effect of employing various concentrations of formic acid and water on phase transition has systematically been studied by XRD, FTIR, and Raman spectroscopy. A considerably higher anatase phase (41%) has been obtained at 800 °C, and 10% anatase composition is retained after annealing the materials at 900 °C for the optimized composition. On comparison, a control sample which has been prepared without formic acid showed that the rutile phase formed at a temperature of 600 °C, FTIR and Raman studies indicated that the formate group favored a bridging (syn–anti or syn–syn) mode of chelation depending on the reaction conditions. It has been concluded that the resulting syn–anti binding hinders cross-linking of the gel network, resulting in a weakened structure and thus causing the anatase to rutile transformation temperature to occur at a lower temperature than with the syn–syn mode of binding where more ordered gel networks are formed.

20 Introduction

Titanium dioxide (TiO_2) is a semiconductor metal oxide that 21 attracts much interest due to its wide range of applications which 22 include photovoltaic cells, gas sensors, pigments, and photo-23 catalysis.¹⁻⁷ It exists as three different polymorphs: rutile 24 25 (tetragonal), anatase (tetragonal), and brookite (orthorhombic).^{8,9} Rutile is thermodynamically stable, while anatase and brookite 26 are metastable, transforming to rutile under calcination, typically 27 600-700 °C.8 Anatase is usually considered to be the most 28 29 photoactive of the three polymorphs for the degradation of organic pollutants.10-12 30

Anatase-to-rutile phase transformation in TiO₂ is an area of 31 both scientific and technological interest.^{6,13} The anatase-to-rutile 32 transformation (ART) is kinetically defined, and the reaction 33 rate is determined by parameters such as particle shape/size,¹⁴ 34 purity,15 source effects,16 atmosphere,17 and reaction conditions.18 35 36 It is agreed that the mechanism for phase transformation of titania is one of nucleation and growth.^{19,20} Anatase nanocrystals 37 coarsen, grow, and then transform to rutile only when a critical 38 size is reached.²¹ Therefore, phase transformation is dominated 39 by effects such as defect concentration,²² grain boundary 40 concentration,²³ and particle packing.²⁴ 41

42 Sol-gel synthesis of TiO_2 is regarded as a relatively straightforward synthesis technique and is thus of great interest 43 and use to researchers.²⁵ Titanium alkoxides are readily 44 hydrolyzed by water due to their susceptibility to nucleophilic 45 attack.²⁶ Rates of hydrolysis and condensation may be controlled 46 by using organic chelating agents such as acetyl acetone,²⁷ 47 alkanolamines,²⁸ diols,²⁹ and acetic acid³⁰ to replace alkoxide 48 groups on the central metal atom. This stability improves control 49

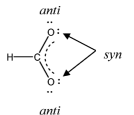


Figure 1. Carboxylate functional group.

over the reaction conditions.³¹ Suresh et al. concluded that the pH of the precursor influences the chelation effect of acetic acid and that the extent of chelation of the acetate group decreases with an increase in pH causing weakened gel structures resulting in decreased ART temperatures.⁶ 54

FTIR and Raman spectroscopy can be used to determine 55 the binding mode of the carboxylate group. The frequency 56 of the asymmetric carboxylate vibration in the IR spectra, 57 $v_{\rm as}(\rm COO^{-})$, and the magnitude of the separation between the 58 carboxylate stretches, $\Delta = v_{as}(COO^{-}) - v_{s}(COO^{-})$, are often 59 used to determine the mode of the carboxylate binding.³² In 60 the bridging coordination, one divalent metal cation is bound 61 to one of the oxygen atoms of the COO⁻ group and another 62 divalent metal cation to the other oxygen, the asymmetric 63 stretch is located at the same position as that of the ionic 64 group.^{32,34} The range 200–210 cm⁻¹ was derived for ionic 65 formates, and in general, the comparison of the Δ value of 66 the respective complex with the Δ value of the sodium salt 67 should be used for the assignment following the guidelines: 68 (i) bidentate chelating coordination, $\Delta(COO^{-})_{\text{formate complex}} \ll$ 69 $\Delta(\text{COO}^-)_{\text{sodium salt}}$; (ii) bidentate bridging carboxylate, 70 $\Delta(\text{COO}^-)_{\text{formate complex}} \leq \Delta(\text{COO}^-)_{\text{sodium salt}}$; (iii) monodentate 71 coordination, $\Delta(COO^{-})_{\text{formate complex}} \gg \Delta(COO^{-})_{\text{sodium salt}}^{32,33,35-38}$ 72

The carboxylate functional group (Figure 1) has four lone 73 pairs of electrons available for coordination to a metal. These 74

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10.1021/jp904358g CCC: \$40.75 © XXXX American Chemical Society

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lone pairs subtend to an angle of 120° and are referred to as 75 76 the syn and anti lone pairs. On the basis of stereoelectronic arguments, it has been suggested that the syn lone pairs are 77 more basic than those in the anti position.³⁹ 78

Previous work studying ART temperatures has been carried 79 out grouping our laboratories where acetic acid was chelated 80 to the titanium alkoxide precursor.⁴⁰ It is widely accepted that 81 acetic acid forms a bridging complex to titanium alkoxides.^{25,41,42} 82 Titanium dioxide sol-gel materials synthesized using titania 83 alkoxides (without chemical additives) typically transform from 84 anatase to rutile at ~ 600 °C. However, to the best of the authors' 85 knowledge, no systematic spectroscopy studies have been carried 86 out to investigate titania carboxylate complexes and how 87 different modes of binding affect the ART temperature. In order 88 89 to determine whether varying ratios of a chelating species alters 90 the ART temperature, the simplest carboxylic acid-formic 91 acid-and titanium isopropoxide in various ratios were used to 92 synthesize a wide range of TiO₂ powders. The formate carboxylate group was chosen because of its versatile coordination 93 behavior.32 The carboxylate coordinations include ionic, mono-94 dentate, bidentate chelating, and bridging.³² From FTIR and 95 Raman spectroscopy, the role of the chelating agent in the 96 synthesis was examined. Ivanda et al. carried out spectroscopic 97 studies on powders synthesized from an esterfication reaction 98 to find bridging of various carboxylates. However, the paper 99 focused mainly on particle size as opposed to relating oligomeric 100 structure and ART temperatures.43 101

102 These are practical, efficient, and useful techniques for gaining 103 information on modes of binding. In order to understand why 104 anatase transforms to rutile at different temperatures, these 105 spectroscopic techniques were employed to investigate how the formate group binds to the titanium. Anatase and rutile 106 percentages and transformation temperatures were examined 107 using XRD. It should also be noted that syn-syn, syn-anti 108 bidentate bridging complexes have not been systematically 109 investigated with relation to ART temperatures. 110

Experimental Section 111

Titanium tetraisopropoxide (TTIP) was added to formic acid 112 under stirring, resulting in a vigorous exothermic reaction 113 producing a powdered suspension. To the resulting suspension 114 water was added in varying ratios to give total volumes in the 115 116 range of 10-50 mL. The molar ratios were varied throughout the experiment: TTIP/formic acid remains constant, 1:2, while 117 the water ratio increases 4, 8, 10, 40, 80, and 100. These samples 118 were labeled F_2W_4 , F_2W_8 , F_2W_{10} , F_2W_{40} , F_2W_{80} , and F_2W_{100} . 119 Similarly, a further 12 samples were prepared where the TTIP/ 120 formic acid ratio was increased to 1:4 and 1:10, respectively, 121 the water ratio was increased to replicate samples $F_2W_4 - F_2W_{100}$. 122 The samples were labeled F_4W_4 , F_4W_8 , F_4W_{10} , F_4W_{40} , F_4W_{80} , 123 and F_4W_{100} when the TTIP/formic acid ratio was 1:4 and for 124 125 the TTIP/formic acid ratio of 1:10 the samples were labeled 126 $F_{10}W_4$, $F_{10}W_8$, $F_{10}W_{10}$, $F_{10}W_{40}$, $F_{10}W_{80}$, and $F_{10}W_{100}$. $F_{control}$ was 127 synthesized without formic acid to compare the chelating effects 128 of the carboxylate group. After the addition of water, a precipitate was formed which was then filtered, washed with 129 deionized water, and dried at 100 °C for 10 h. Each powder 130 was then calcined at temperatures ranging from 300-1000 °C. 131 XRD patterns of the calcined powders were obtained with a 132 Siemens D500 X-ray powder diffractometer in the diffraction 133 angle range $2\Theta = 20-80^\circ$ using Cu K α radiation. Anatase/ 134 rutile percentages were calculated from the resulting diffracto-135 grams using the Spurr equation:⁴⁴ 136

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$$\%_{\text{Rutile}} = \frac{1}{1 + 0.8[I_{\text{A}}(101)/I_{\text{R}}(110)]}$$
(1)

Where I_A is the intensity of (101) peak and I_R is the intensity 137 of (110) peak. 138

Infrared and Raman spectroscopy were employed to determine carboxylate binding modes on each powder before or xerogel calcination.

All infrared spectra were obtained on a Perkin-Elmer GX FT-142 IR and recorded as a KBr disk (1:10 sample/KBr). An ISA 143 Labram was used to record Raman spectra of the powdered 144 samples with an argon ion laser (514.5 nm) as excitation source. 145

FESEM images were obtained from a Hitachi SU-70 at an 146 operating voltage of 5.0 kV. 147

Methylene blue is an accepted model organic pollutant for 148 photocatalytic degradation studies and is used as an industrial 149 standard (Japanese standard, JIS R 1703-2:2007); as such, it 150 was used to demonstrate the photocatalytic efficiency of the 151 synthesized TiO₂ powders. Rhodamine 6G (5 \times 10⁻⁶ M) was 152 used for the comparison of results. In a typical experiment, 153 crystalline TiO_2 (60 mg) was added to methylene blue solution 154 (50 mL, 1 \times 10 ⁻⁵ M) and placed in a Q-sun solar simulator⁴⁵ 155 with continuous stirring, 5 mL aliquots were withdrawn at timed 156 intervals and the visible absorption spectrum was measured 157 using a Perkin-Elmer Lambda 900 UV-vis spectrometer. 158

Results and Discussion

In order to investigate the influence of chelation on ART, 160 the effect of various concentrations of water and formic acid 161 on the titania precursor were investigated. 162

Effect of Water. X-ray Diffraction. XRD was employed to 163 determine the phase analysis of each powder calcined at 164 increasing temperatures. The percentage of anatase in the 165 calcined sample is shown in Figure 2. All samples were 100% 166 F2 anatase at temperatures 300-500 °C. From Figure 2 it is 167 apparent that for each series of powders $(F_2W_4-F_2W_{100})$, 168 F_4W_4 - F_4W_{100} , and $F_{10}W_4$ - $F_{10}W_{100}$), where the TTIP/FA ratio 169 remains constant, that the increase in water promotes the 170 formation of rutile. For the powders $F_2W_4 - F_2W_{100}$, F_2W_4 has 171 the lowest water ratio and it is the only powder with anatase at 172 700 °C, but the remaining powders have all underwent complete 173 phase transformation to a more thermodynamically stable rutile. 174

The formic acid ratio was increased for powders 175 $F_4W_4 - F_4W_{100}$, causing improved chelation. This is reflected 176 throughout the six powders, as most retain anatase at 700 °C. 177 Anatase is predominant for F_4W_4 at 700 °C (86%), but the 178 increase in water along the series causes increased rutile 179 formation for the remaining powders. At 800 °C anatase is still 180 dominant for the powder F_4W_4 (61%) and is even present in 181 F_4W_8 (5%); however, the other powders in the series are rutile. 182

From these results it is clear that an increase in the amount 183 of water used for hydrolysis has an adverse effect on ART 184 temperatures, resulting in the lowering of the ART. This increase 185 in water reduces the acidity of the sol resulting in a decrease in 186 the chelation effect of the formate group.^{26,41,42,46} This decrease 187 in chelation results in a weakened gel network and consequently 188 a lowering of ART temperature was observed.⁴⁷ Sahni et al. 189 reported that increasing water content causes increased hydroly-190 sis which results in the formation of larger particles that 191 thermodynamically favor phase transformation to rutile.⁴⁸ 192

For powders $F_{10}W_4 - F_{10}W_{100}$, the sample with the lowest 193 water ratio, $F_{10}W_4$, is one of only two samples where anatase 194 is present at 800 °C. The other is $F_{10}W_{40}$, and the reason why 195

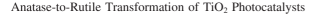
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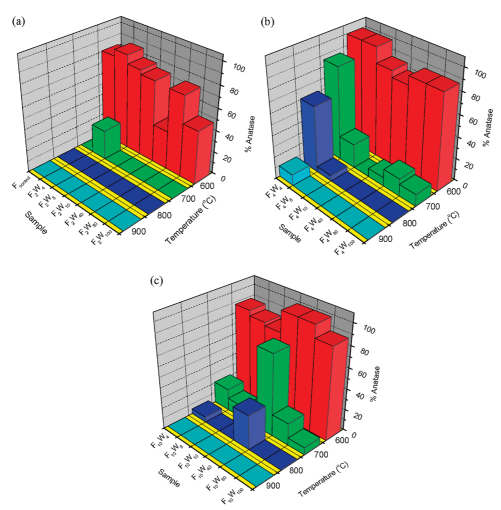


Figure 2. Percentage of anatase in the calcined TiO₂ samples, determined by XRD for materials heated to different temperatures (a) $F_2W_4 - F_2W_{100}$, (b) $F_4W_4 - F_4W_{100}$, and (c) $F_{10}W_4 - F_{10}W_{100}$.

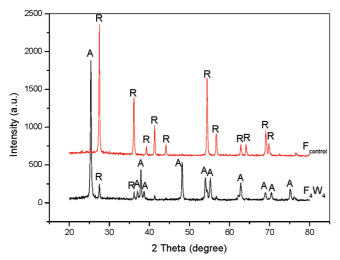


Figure 3. XRD of F_{control} and F₄W₄ at 700 °C. A, anatase; R, rutile.

anatase is present in $F_{10}W_{40}$ is unclear as it has an increased amount of water, whereas powders $F_{10}W_8$, $F_{10}W_{10}$, $F_{10}W_{80}$, and $F_{10}W_{100}$ follow the theory that increased amounts of water cause a reduction in ART temperatures.

From Figure 2, F_4W_4 was found to maintain anatase at temperatures as high as 900 °C. All powders synthesized (bar $F_{10}W_{40}$) showed that as water ratio is increased rutile forms at lower temperatures.

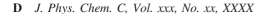
As the carboxylate group may coordinate to Ti in a number 204 of different arrangements^{26,32,33,41,42,46} spectroscopic (Raman and 205 IR) studies were carried out on all precalcined samples in order 206 to determine if an increase in water influences the way that the 207 carboxylate group binds to Ti, which in turn affects the anatase 208 to rutile transformation temperature. The morphology of the 209 calcined powders were analyzed using FESEM microscopy and 210 were found to be highly aggregated with an average size of 211 100–150 nm (Supporting Information 1). 212

Infrared Spectroscopy. Figure 4 shows the IR spectra of 213 F4 powdered samples F_2W_4 and F_2W_8 ($F_2W_8-F_2W_{100}$ give near 214 identical spectra). At \sim 450 cm⁻¹ there is a broad peak due 215 to the formation of Ti-O bonds. Peaks at 1350 and 1550 216 cm⁻¹ represent $v(COO^{-})_{sym}$ and $v(COO^{-})_{asym}$, respectively. 217 Zalenak et al. determined the mode of bonding of the 218 carboxylate group to the metal atom by calculating Δ , where 219 $\Delta = v_{\rm as}(\rm COO^-) - v_s(\rm COO^-).^{32}$ From the spectra obtained 220 for F_2W_4 and F_2W_8 (Figure 4), $\Delta = \sim 200 \text{ cm}^{-1}$. This value 221 is consistent with the literature value for ionic formate 222 $(\text{HCOO}^{-}) = \sim 201 \text{ cm}^{-1}.^{33}$ 223

As has been reported elsewhere, the bidentate bridging carboxylate exists when $\Delta(\text{COO}^-)_{\text{formate complex}} \leq 225$ $\Delta(\text{COO}^-)_{\text{sodium salt.}}^{33,35-38}$ Therefore, it is proposed that the 226 formate group binds to the Ti center in bidentate bridging mode 227 such as syn-syn or syn-anti (Figure 5). 228 F5

It has been reported that metal alkoxo-acetates are formed 229 by the reaction of acetic acid (AcOH) with metal alkoxide, 230

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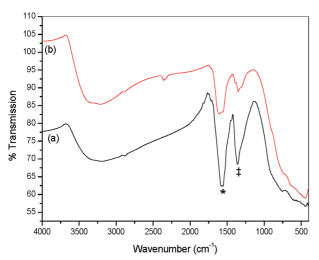


Figure 4. IR spectra of TiO₂ precursor powders (a) F_2W_4 and (b) F_2W_8 , where $* = v_{asym}(COO^-)$ and $\ddagger = v_{sym}(COO^-)$.

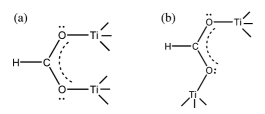


Figure 5. Bidentate bridging modes of the formate group and TTIP (a) syn-syn and (b) syn-anti.

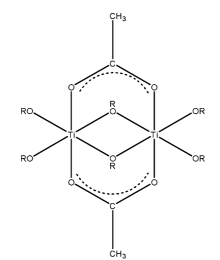


Figure 6. Bridging acetate ligands attached to TTIP.

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in which OPrⁱ (OR) groups on the central titanium atom are
preferentially hydrolyzed, whereas bridging acetate ligands
remain bonded to titanium throughout much of the condensation process (Figure 6).^{25,42}

Since the bridging acetate ligands are not hydrolyzed, the 235 chelated ligands effectively alter the condensation pathway 236 237 toward promoting the formation of linear polymers composed of edge-sharing octahedra.42,49 The addition of excess water 238 destabilizes the system by altering the highly cross-linked 239 240 network structure. The gel having polymeric chains with little branching and cross-linking, as well as a smaller void region, 241 are structurally weak and thus collapse rapidly on calcination 242 forming rutile at lower temperatures.⁴⁷ 243

This can clearly be seen from both the rutile percentage results (Figure 2) and also from the IR spectra (Figure 7) where bands corresponding to the chelated formate group become weaker.

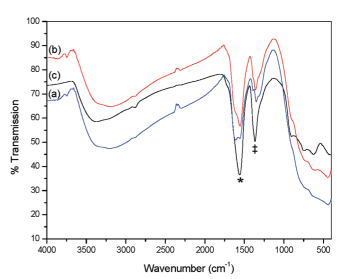


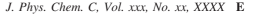
Figure 7. IR spectra of TiO₂ precursor powders (a) F_4W_4 , (b) F_4W_8 , and (c) F_4W_{10} , where $* = v_{asym}(COO^-)$ and $\ddagger = v_{sym}(COO^-)$.

From the IR spectra of samples $F_4W_4 - F_4W_{10}$ (Figure 7), it can 247 be seen that there is a strong OH peak at $\sim 2800-3500$ cm⁻¹. 248 As the ratio of water is increased in each series $(F_2W_4 - F_2W_{100})$, 249 $F_4W_4-F_4W_{100}$, $F_{10}W_4-F_{10}W_{100}$) it can be seen that the intensity 250 of both $v_{sym}(COO)$ and $v_{asym}(COO)$ decrease in relation to both 251 the Ti–O and the OH peaks. F_4W_4 shows the strongest COO 252 stretches and the weakest Ti-O and OH signal when compared 253 with the other samples. It is believed that this is due to an 254 increase in the molar ratio of water which alters the pH of the 255 system as well as increasing hydrolysis and weakening the gel 256 network.⁶ This, as has been reported previously, weakens the 257 chelation of the carboxylate group, which will cause weakened 258 COO stretches, facilitating increased hydrolysis, therefore 259 increasing OH stretches and causing a reduction in the anatase 260 to rutile transformation temperature.6,47,48 261

It can also be seen from the IR spectra (Figures 4 and 7) that 262 for samples F₂W₈ and F₄W₁₀ two asymmetric (1600 and 1550 263 cm^{-1}) and two symmetric (1382 and 1340 cm^{-1}) carboxylate 264 stretches were observed. For sample F_4W_4 one asymmetric (1550 265 cm⁻¹) and one symmetric (1362 cm⁻¹) carboxylate stretch were 266 observed in the IR spectra. The spectra of sample F₄W₈ showed 267 that secondary asymmetric and symmetric stretches were 268 beginning to form. The presence of two carboxylate stretches 269 indicates the presence of two different modes of carboxylate 270 binding.³² The frequency of asymmetric and symmetric vibra-271 tions depends on the electronic charge density of C-O bonds 272 and C-O bond lengths; the higher is the frequency of the 273 asymmetric vibration, and the lower is the frequency of the 274 symmetric vibration. Hence for compounds F₂W₄, F₄W₄, F₄W₈, 275 and $F_{10}W_{40}$ the asymmetric vibration at 1550 cm⁻¹ pertains to 276 the symmetric vibration at 1350 cm⁻¹. Similarly, the stretches 277 at 1600 and 1380 cm⁻¹ are related. The respective experimental 278 values are $\Delta_{exp} = {\sim}210~\text{cm}^{-1}$ for samples $F_2W_4,~\tilde{F}_4W_4,$ and 279 F_4W_{10} , and $\Delta_{exp} = \sim 220 \text{ cm}^{-1}$ for the remaining samples. Both 280 Δ_{exp} are similar and indicate the bridging chelation. However, 281 the formation of secondary peaks indicates a different binding 282 mode. As stated above, the carboxylate functional group has 283 two lone pairs of electrons on each oxygen atom available for 284 binding, the syn lone pair and the anti lone pair. It has been 285 suggested that the syn lone pair is more basic than those in the 286 anti position.³⁹ It may be possible that a syn-anti (Figure 5b) 287 mode of binding occurs when the water ratio is increased due 288

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Anatase-to-Rutile Transformation of TiO₂ Photocatalysts



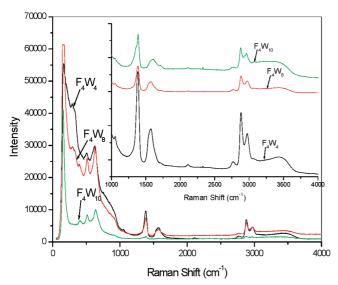


Figure 8. Raman spectra of TiO_2 precursor powders of $F_4W_4 - F_4W_{10}$.

to the altering pH and the increased hydrolysis of the system.This would explain the appearance of secondary peaks in theIR spectra.

Zelenak et al. observed singlet peaks for both the v_{as} and v_{s} 292 (COO⁻) stretches for bridging complexes that follow syn-syn 293 294 binding and they observed doublet peaks at similar wavenumbers for syn-anti binding with zinc carboxylate complexes,³² 295 while Ishioka et al. reported similar values of separation for 296 syn-anti bridge in zinc(II) acetate.⁵⁰ The result of syn-anti 297 bridging may produce a polymeric network with little branching 298 and cross-linking that is structurally weak, therefore, forming 299 rutile at lower calcination temperatures than that of a typical 300 syn-syn mode of bidentate bridging. 301

Raman Spectroscopy. Raman spectroscopy was employed
 as a secondary technique to IR in order to confirm the above
 results.

F8

Figure 8 shows the Raman spectra of the precalcined TiO₂ 305 powders. Although the powders have not been calcined, the 306 Raman spectra display clear signs of the anatase phase four-307 308 peak pattern with peaks at 160, 405, 515, and 635 cm^{-1} for 309 powders F_4W_{10} - F_4W_{100} . However, for the powders F_4W_4 and F_4W_8 (like with IR spectra) the appearance of a peak at ~290 310 cm⁻¹ indicates that the Ti–O structure is different than the other 311 samples and contains a pattern similar to that of an anatase/ 312 rutile mixture. The formation of rutile-like structures during the 313 course of crystallization of titania hydrolysate into anatase has 314 been confirmed by several research groups.⁵¹⁻⁵³ It has been 315 suggested that the structures which provide the anatase and rutile 316 Raman spectral patterns disappear just before the crystallization 317 into anatase.⁵³ It is apparent that the presence of the anatase/ 318 rutile-like structure for samples F_4W_4 and F_4W_8 cause an 319 increase in the ART temperature. From Figure 8 it can also be 320 seen that the presence of intense peaks at 1393, 1580, 2890, 321 322 and 2980 cm⁻¹ are only present in samples F₄W₄ and F₄W₈. In 323 order to investigate further, all Raman spectra were repeated but scans were only carried out in the region $800-4000 \text{ cm}^{-1}$ 324 325 (the organic region) to further confirm binding modes of the formate group. 326

Figure 8 (inset) shows the Raman spectra of the formate group binding with the titanium. Peaks from $\sim 2800-3500$ cm⁻¹ are due to OH stretches.³³ As seen with the IR spectra, the intensity of the OH peak increases in comparison with the COO⁻ stretches (1392 and 1567 cm⁻¹) in F₄W₄ and F₄W₈ when compared with

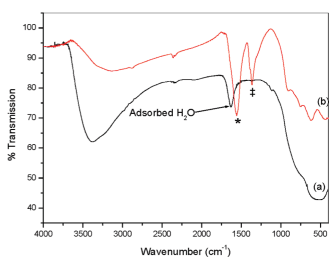


Figure 9. IR spectrum of TiO₂ precursor powders (a) $F_{control}$ and (b) F_4W_4 , where $* = v_{asym}(COO^-)$ and $\ddagger = v_{sym}(COO^-)$.

 $F_4W_{10}-F_4W_{100}$. The appearance of secondary peaks also appears beside the main COO⁻ peaks (1392 and 1567 cm⁻¹) at 1370 and 1720 cm⁻¹. The peak at 1567 cm⁻¹ is also shifted to a higher wavenumber. The formation of secondary peaks in the Raman spectra are for the same samples as those of the IR which again indicates the presence of an alternative mode of binding such as syn-anti, as was proposed previously (Figure 5). 332

Effect of Chelating Agent. Sample F_{control} was synthesized 339 using water only to determine what affect formic acid had on 340 the structure of the Ti–O network both before and after 341 calcination. 342

X-ray Diffraction. Without the presence of a chelating agent 343 (F_{control}), rutile begins to form at a temperature as low as 344 600 °C (20%) and total transformation has occurred at 700 °C. 345 Samples F₂W₁₀, F₂W₄₀, F₂W₈₀, and F₂W₁₀₀ have a higher rutile 346 content at 600 °C. This may be due to the chelating agent having 347 an adverse effect on the initial TiO₂ structure whereby a 348 syn-anti bridging mode is dominant throughout the structure 349 thus forming a structure without cross-linking that upon 350 calcination, forms a larger percentage of rutile at 600 °C. 351

Infrared Spectroscopy. In Figure 9 there is an OH stretch 352 $(2800-3600 \text{ cm}^{-1})$, a Ti-O stretch $(400-1000 \text{ cm}^{-1})$, and also 353 a signal at 1610 cm⁻¹ due to the bending vibrations of adsorbed 354 water. There is a clear difference in the IR spectra of F_{control} 355 and F_4W_4 . This was expected and is due to the carboxylate 356 group-Ti bridging structure. Also in the region 400-1000 357 cm^{-1} , F_4W_4 gives more defined peaks as opposed to the broad 358 peak given by $F_{control}$. This is due to a more ordered Ti–O 359 framework.26,31,43 360

Raman Spectroscopy. The Raman spectrum of F_{control} gives 361 no peaks of distinction. Showing that without the presence of 362 formic acid, Ti-O atoms randomly arrange as opposed to the 363 more ordered structure shown in Figure 8 where formic acid 364 was employed as a chelating agent. Comparing the Raman 365 spectra of F_{control} with Figure 9, where the samples were chelated, 366 it becomes clear that the presence of the formic acid as a 367 chelating agent enables the metal-oxygen atoms to form a 368 defined, crystallinelike structure which is apparent in Figure 8 369 $(0-1000 \text{ cm}^{-1})$. The Raman spectra of F_{control} , as expected also 370 lacks the presence of the bridging peaks present in Figure 8 at 371 1390 and 1570 cm^{-1} . It is the presence of this CO-Ti bridge 372 that allows a controlled arrangement of the Ti-O atoms. 373 Without the bridge there is uncontrolled hydrolysis leading to 374 a random arrangement of Ti-O atoms. The Raman spectra of 375

F9

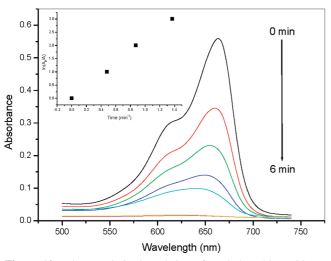


Figure 10. Photocatalytic degradation of methylene blue with F7 (700 °C).

TABLE 1: Reaction Rate Constants (± 0.01 , k/\min^{-1}) for the Degradation of Methylene Blue

sample	600 °C	700 °C	800 °C
F _{control}	0.12	0.12	0.05
F_2W_4	0.17	0.20	0.17
F_4W_4	0.13	0.45	0.38
$F_{10}W_4$	0.26	0.13	0.08

 F_{control} compares favorably to the IR spectrum of the same sample shown in Figure 9.

Photocatalytic Studies. Photocatalytic studies were carried out on selected powders ($F_{control}$, F_2W_4 , F_4W_4 , and $F_{10}W_4$) at calcination temperatures (600, 700, and 800 °C) and were compared with the commercial photocatalyst, Degussa P25. Powder F_4W_4 calcined at 700 °C was found to be the most photocatalytically active (Figure 10). The methylene blue was completely degraded after 6 min.

F10

T1

Improved photocatalytic activity has been previously found 385 386 with anatase/rutile interactions, due to improved electron-hole separation.⁵⁴⁻⁵⁶ A mixture of both phases has given rise to the 387 most efficient photocatalyst out of the powders synthesized. It 388 has been suggested that an intimate contact between anatase 389 and rutile phases may enhance the separation of photogenerated 390 391 electrons and holes resulting in excellent photocatalytic efficiency.⁵⁷ It is believed that the anatase/rutile mixture present 392 in Degussa P25 is one of the reasons why it is the one of the 393 most investigated photocatalysts.56 First-order degradation plots 394 of powders $F_{control}$, F_2W_4 , F_4W_4 (Figure 10 inset), and $F_{10}W_4$ at 395 396 calcination temperatures 600, 700, and 800 °C were used to calculate the reaction rate constant, $k \pmod{1}$ (Table 1). 397

Sample F₄W₄ calcined at 700 °C had the largest rate constant 398 at 0.45 min⁻¹, the rate constant for Degussa P25 was found to 399 be 0.29 min^{-1} for the same reaction conditions. Sample F_4W_4 400 calcined at 700 °C consists of 87% anatase and 13% rutile to 401 402 give an ideal mixture for photocatalytic efficiency for the 403 degradation of methylene blue as a model pollutant. Sample F₂W₄ has an identical anatase/rutile mixture at 600 °C but is 404 405 not as photoactive as F₄W₄. At 700 °C F₂W₄ consists of mainly rutile (72%) which results in a reduction of photocatalytic 406 activity. At 600 and 800 °C, F4W4 has 0% and 32% rutile, 407 respectively. This reduces the photocatalytic activity of F₄W₄. 408 F_{control}, the sample prepared without any chelating agent was 409 410 the poorest photocatalyic performer, even at 600 °C where it consisted of an anatase/rutile mixture (90:10), giving a reaction 411

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rate of 0.12 min⁻¹. As shown through IR and Raman spectros-412 copy, the absence of a chelating agent causes complete disorder 413 among the Ti-O bonds upon hydrolysis resulting in an 414 unorganized network of TiO₂ particles when compared with 415 those where a chelating agent was present. $F_{10}W_4$ consisted of 416 77% rutile at 600 °C and 95% rutile at 700 °C. As reported 417 previously,⁵⁴⁻⁵⁷ a mixture of anatase and rutile has greater 418 photoactivity than either phase alone. Photocatalytic studies were 419 repeated for the same samples using rhodamine 6G as a model 420 organic pollutant and consistent results were obtained (Sup-421 porting Information 2). 422

Conclusions

ART in a formic acid modified titania material has been 424 studied using XRD, FTIR, and Raman spectroscopy. Through 425 Raman and IR it was possible to determine the mode of binding 426 of the chelating agent, formic acid, to the titanium precursor 427 with the equation $\Delta = v_{as}(COO^{-}) - v_s(COO^{-})$. A value for 428 $v_{\rm as}(\rm COO^{-}) - v_{\rm s}(\rm COO^{-})$ of 210 cm⁻¹ indicated that bidentate 429 bridging is the mode of binding for the samples. However, for 430 samples with increased water concentrations, spectroscopy 431 results showed doublet peaks indicating alternate modes of 432 bridged binding. It was postulated that for these samples 433 syn-anti binding was occurring as well as syn-syn binding. It 434 is believed that the resulting syn-anti binding hinders cross-435 linking of the oligomer network, resulting in a weakened 436 structure and thus causing the anatase to rutile transformation 437 temperature to occur at lower temperatures than with the 438 syn-syn mode of binding where more ordered oligomer 439 networks are believed to be formed. 440

Photocatalytic studies showed that the formic acid modified sample (calcined at 700 °C) with an anatase/rutile mixture of 86:14 was more effective for the degradation of methylene blue than the commercial titania sample Degussa P25, showing that an anatase/rutile mixture is more effective than either phase alone, which is consistent with previous literature results. 440

Acknowledgment. The authors thank the Environmental447Protection Agency for its financial support. They also thank448Emer Ryan for FESEM images.449

Supporting Information Available:FESEM image of450calcined powders and photocatalytic testing data for rhodamine4514516G. This material is available free of charge via the Internet at452http://pubs.acs.org.453

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