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# Highly Visible Light Active TiO2-xNx Heterojunction Photocatalysts

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## Highly Visible Light Active $TiO_{2-x}N_x$ Heterojunction Photocatalysts<sup>†</sup>

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Nitrogen doped anatase-rutile heterojunctions are successfully synthesized through an ethylenediaminetetraacetic acid (EDTA) modified sol-gel process. An FT-IR study of EDTA modified TiO<sub>2</sub> gel confirms the existence of an ionic intermediate (as indicated by a  $\Delta \nu$  value of 233 cm<sup>-1</sup>). Differential scanning calorimetry (DSC), X-ray diffraction (XRD), and Raman spectroscopy are employed to study the phase evolution, phase purity, and crystallite size of samples. Formations of O-Ti-N and N-Ti-N bonds in calcined samples are confirmed using XPS and FT-IR spectroscopy. All EDTA modified samples show significantly higher visible light photocatalytic activity than the unmodified sample. The most active nitrogen doped heterojunction obtained at 400 °C exhibits 9-fold visible light activity in comparison to the standard photocatalyst Degussa P-25. It is proposed that the photo excited electrons (from the visible midgap level) are effectively transferred from the conduction band of anatase to that of rutile causing effective electron-hole separation, which is responsible for the higher visible light activity and lower photoluminescence (PL) intensity.

#### 1. Introduction

Unique properties of nanosized catalysts such as quan-21 tum confinement and high surface to volume ratio make 22 23 them highly efficient. Among various catalysts, titania nanomaterials find wide applications in the field of solar 24 cells,<sup>1</sup> organic synthesis,<sup>2</sup> water and air purification,<sup>3</sup> cancer 25 therapy,<sup>4</sup> cathodic corrosion protection, and self-cleaning 26 antibacterial materials.<sup>5,6</sup> High redox potential, chemical 27 stability, inexpensiveness, and non toxicity of titania made 28 it superior to other semiconductor photocatalysts in its 29 class. Among the three polymorphs, it is generally agreed 30 that the anatase phase has the highest photoactivity fol-31 lowed by rutile and brookite. The efficiency of titania 32 photocatalysts strongly depends on several factors, includ-33 ing phase purity, surface area, crystallite size, amount and 34 nature of dopants, method of preparation, and anatase-35 rutile ratio.7-9 36

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The wide band gap (3.2 eV) of the anatase titania 37 confines its application to UV light ( $\lambda \leq 387.5$  nm) 38 activation. Even though the high temperature stable rutile 39 phase can absorb visible light (up to  $\sim$ 412 nm), its activity 40 is limited as a result of low surface area, low redox potential, 41 and faster electron hole recombination rate.<sup>10</sup> The deve-42 lopment of a highly visible light active catalyst is necessary 43 to effectively exploit sunlight or light from artificial sources. 44 Various methods such as metal or non-metal doping, using 45 reduced forms of  $TiO_X$  (X < 2) photocatalyst, dye sensiti-46 zation, and optimization of phase composition has been 47 investigated by previous researchers.<sup>11-13</sup> 48

Titania doped with main group elements has attracted 49 great interest after the report of nitrogen and other anion 50 (S, C, and F) doped visible-light active (VLA) titania cata-51 lyst by Asahi and co-workers.<sup>14</sup> These second-generation 52 TiO<sub>2</sub> materials were photoactive over the UV and visible-53 light region. Controversy still remains about the nature of 54 the dopant species and electronic structure of nitrogen 55 doped titania, which is highly dependent on the method of 56 preparation. Some researchers proposed the existence of 57 surface adsorbed NO<sub>X</sub> and NH<sub>X</sub>, while others proposed the 58 presence of lattice nitrogen. Recently, Valentine et al. 59 demonstrated the electronic structure of nitrogen doped 60

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<sup>&</sup>lt;sup>†</sup> This paper is dedicated to Professor John M. Kelly, C. Chem. FRSC on the occasion of completing his 37 years of teaching and research (celebration of chemistry) at the Trinity College Dublin, Ireland.

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titania synthesized through various routes.<sup>15</sup> Density 61 functional theory (DFT) calculations show that substitu-62 tional nitrogen atoms create an energy level just above the 63 valence band, where as interstitial nitrogen energy levels 64 lie higher in the gap. Calculations also showed that nitro-65 gen doping leads to an ample lowering of oxygen vacancy 66 formation energy. Photoelectrochemical studies of nitro-67 gen doped titania revealed the formation of additional 68 energy states positioned energetically 1.3 eV broad range 69 just below the conduction band edge. These energy levels 70 cause higher loss of visible light excited charge carriers as 71 a result of recombination.<sup>16</sup> Increasing the doping density 72 also enhances the electron-hole recombination as a result 73 of the reduced band gap. These main drawbacks of nitrogen 74 doped titania limit its visible light activity and applications. 75 For efficient visible light photocatalysis, coupling of nitro-76 77 gen doped titania with the electron hole separating agents are necessary. 78

One way to attain high activity is proper designing of 79 heterojunctions with smaller band gap semiconductors, 80 such as rutile. A mixture of anatase and rutile can often 81 82 outperform the photoactivity of pure anatase or rutile. Recent studies explained the visible light induced produc-83 tion of charges in smaller band gap rutile phase and its 84 transfer to the trapping states of anatase phase.<sup>17</sup> Since 85 the conduction band level of rutile is below that of anatase, 86 photo generated electrons from the conduction band of 87 anatase can be effectively transferred to that of rutile 88 leading to slow electron hole recombination.<sup>7,18</sup> These 89 two key factors are reported to be responsible for the 90 superior photocatalytic activity of Degussa P25, which 91 has an anatase-rutile ratio of 70/30.<sup>19</sup> Most of the synthesis 92 methods of titania gives anatase rutile mixtures only above 93 600 °C resulting in a deviation from nanoregime and a 94 subsequent decrease in photocatalytic activity. Nitrogen 95 doping also results in the formation of high temperature 96 stable anatase phase titania.<sup>20-22</sup> Under these circum-97 stances, it is necessary to develop a low temperature 98 method for the synthesis of nitrogen doped anatase-rutile 99 nano heterojunctions. This has been achieved through an 100 ethylenediaminetetraacetic acid (EDTA) modified sol-101 gel process, which proceeds through an ionic intermediate 102 [Ti (H<sub>2</sub>O) (edta)]. This method proceeds through a solu-103 tion phase without the formation of polymeric chains, 104 which can ensure better homogeneity and effective doping 105 than conventional sol-gel processes. Both low temperature 106

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crystallizations of anatase-rutile heterojunctions and nitro-107 gen doping were achieved using EDTA. Nishide et al. 108 studied the crystal structure and optical properties of 109 TiO<sub>2</sub> materials prepared from Ti-EDTA complexes.<sup>23</sup> High 110 photocatalytic activities of anatase-brookite and anatase-111 carbon nanotubes heterojunctions have also been re-112 ported.<sup>24,25</sup> However, to the best of our knowledge, there 113 is no systematic study available on a highly visible light 114 active  $TiO_{2-x}N_x$  anatase-rutile heterojunction photoca-115 talysts. The current study focuses on the low temperature 116 crystallization, mechanism of formation, and the photo-117 catalytic activity of  $TiO_{2-x}N_x$  heterojunctions. 118

### 2. Experimental Methods

2.1. Synthesis of N-Doped Titania Heterojunctions. All re-120 agents were used without further purification. Titanium iso-121 propoxide (Aldrich 97%) and EDTA (Aldrich 99.99%) was 122 used as the titania precursor and modifier respectively. In a 123 typical synthesis for 1:1 EDTA modified sample, titanium iso-124 propoxide (9 mL) was dissolved in isopropanol (23 mL). This 125 solution was mixed with glacial acetic acid (1.7 mL) and EDTA 126 (8.8 g). The precipitate obtained was then mixed with deionized 127 water (55 mL) and stirred for 2 h to form a sol. The sol obtained 128 was then heated in an oven at 80 °C to form a solution, which on 129 further heating converts to a gel. Xerogel thus obtained after 24 h 130 was calcined at 400, 500, 600, 700, 800, and 900 °C for 2 h at a 131 heating rate of 10 °C/min. Various molar ratios of TTIP/IPA/ 132 ACOH/H2O/EDTA were used for the synthesis of different 133 samples (Supporting Information 15). All nitrogen doped hetero-134 junctions were prepared by successively replacing 0.5 molar ratios 135 of acetic acid by EDTA (identified as 0.5 ED-TiO<sub>2</sub>, 1.0 ED-TiO<sub>2</sub>, 136 1.5 ED-TiO<sub>2</sub>, and 2.0 ED-TiO<sub>2</sub> respectively after calcination). 137 Ratios between other reagents were maintained as constant in all 138 synthesis. The control sample without EDTA was also prepared 139 using the same synthesis method. Molar ratio between titanium 140 isopropoxide, acetic acid, isopropanol, and water was 1:2:10:100 141 for the synthesis of the control sample. Degussa P-25 was used as 142 the standard TiO<sub>2</sub> sample for comparison. 143

2.2. Characterization Techniques. The FTIR spectra of xero-144 gel dried at 100 °C and calcined at different temperatures were 145 recorded using a Perkin-Elmer GX-FTIR spectrometer in the 146 range 4000-400 cm<sup>-1</sup>. Transparent pellets were prepared using 147 a 4 mm dye after mixing samples with KBr. Rheometric 148 Scientific DSC QC and Shimadzu DTG-60 instruments were 149 used for differential scanning calorimetry (DSC) and differen-150 tial thermal analysis (DTA) respectively. In both cases, about 151 5 mg of dried gel was heated from room temperature (25 °C) to 152 600 °C at a constant heating rate of 10 °C/min. The crystal 153 phases of calcined samples were analyzed by X-ray diffraction 154 using a Siemens D500 X-ray diffractometer ( $2\theta = 10-70^\circ$ ) 155 working with Cu–K $\alpha$  radiation ( $\lambda = 0.15418$  nm). The Spurr 156 equation (eq 1) was employed for the precise calculation of the 157 amount of rutile in the sample. 158

$$F_R = \frac{1}{1 + 0.8[I_A(101)/I_R(110)]} \tag{1}$$

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159 Where  $F_R$  is the mass fraction of rutile,  $I_A(1 \ 0 \ 1)$  and  $I_R(1 \ 1 \ 0)$  are 160 the integrated main peak intensities of anatase and rutile, respectively. Crystallite sizes of anatase and rutile in calcined 161 162 samples were calculated using the Scherrer equation (eq 2).

$$\Phi = \frac{k\lambda}{\beta\cos\theta} \tag{2}$$

Where  $\Phi$  is the crystallite size, k is the shape factor,  $\lambda$  is the X-ray 163 radiation wavelength, and  $\beta$  the full line width at half-maximum 164 165 height of the main intensity peak after subtraction of the equip-166 ment line broadening. Raman spectra of all samples were recor-167 ded at room temperature with a Dilor ISA Labram 1 B micro-Raman system equipped with a 514 nm Ar<sup>+</sup> ion laser (Laser 168 Physics Reliant 150 Select Multi-Line). Measurements were 169 carried out on powder samples at a laser power of 3 mW to 170 171 avoid excessive heating.

X-ray photoelectron spectroscopy (XPS) analyses were per-172 173 formed on a Thermo VG Scientific Sigma Probe spectrometer using monochromatic Al-Ka radiation (photon energy 1486.6 174 eV). For Ti 2p high-resolution spectra pass energy of 20 eV and a 175 0.1 eV step size were used. A pass energy of 50 eV and a step size 176 of 0.2 eV were used for N 1s high-resolution spectra. Charge 177 178 compensation was achieved by using a low-energy electron flood gun. Quantitative surface chemical analyses were calcu-179 lated from the high-resolution core level spectra, following the 180 181 removal of a nonlinear Shirley background. Absorbance spectra 182 of the samples were recorded in the diffuse reflectance mode (with an integrated sphere attachment) using a Perkin-Elmer 183 Lambda 900 UV/vis/NIR spectrometer, using BaSO<sub>4</sub> as the 184 reference sample (in the range of 300-600 nm). Sample pellets 185 were prepared using a 4 mm die after thoroughly mixing the 186 powder samples with KBr. The band gaps were calculated by 187 extrapolating the lower wavelength cutoff region. Samples for 188 photoluminescence studies were prepared by dispersing as pre-189 pared powders (0.01 g in 100 mL) in deionized water. All samples 190 191 were sonicated for 30 min before analysis. Room temperature 192 emission spectra of samples were obtained from Perkin-Elmer 193 Luminescence Spectrometer (LS-55) at an excitation wavelength of 260 nm in the range 300 to 800 nm. 194

Nitrogen adsorption and desorption isotherms were collected 195 at liquid nitrogen temperature using a Quantachrome 2000e 196 surface area analyzer. All samples were degassed at 300 °C for 197 2 h under vacuum prior to the analysis. The specific surface area 198 was calculated using the linear portion of the Brunauer-199 Emmett-Teller (BET) model ( $P/P_0 = 0.05-0.2$ ). Desorption 200 branch of the Barret-Joyner-Halenda (BJH) model was em-201 202 ployed for the precise determination of pore diameter and volumes. For transmission electron microscopy (TEM) ana-203 204 lysis, powder samples were suspended in water and sonicated for 205 30 min before being dispersed onto Formvar-coated copper grids. The particle-loaded grids were studied under JEOL JEM-206 207 2100 transmission electron microscope operating at an accelerating voltage of 200 kV. 208

2.3. Photocatalysis Study. Decomposition of methylene blue 209 was performed to evaluate the photocatalytic activities of samples. 210 The catalyst (0.06 g) was dispersed in an aqueous solution of 211 methylene blue (50 mL  $1 \times 10^{-5}$  M) placed in a glass vessel 212 (100 mL). The vessel was placed in the dark for 30 min and 213 irradiated with visible light  $(0.68 \text{ W/m}^2)$  using Q-Sun solar simu-214 lator and a primary blue filter (450 nm) (Supporting Informa-215 tion 13). The temperature of the suspension was maintained 216 217 at 25 °C with the help of an air cooler (thermostat attached) 218 connected to the solar simulator. Degradation of methylene blue



Figure 1. FT-IR Spectrum of 80 °C dried titania precursor (a) acetic acid modified and (b) EDTA modified.

was monitored by taking aliquots at equal time intervals of 1 h. 219 These aliquots were centrifuged, and absorption spectra of the 220 samples were recorded using Perkin-Elmer Lambda 900 UV/vis/ 221 NIR spectrometer. Similar experiments were carried out for 222 samples calcined at different temperatures. The rate of degrada-223 tion was assumed to obey pseudo-first order kinetics, given 224 that natural logarithmic plots were linear. The rate constant for 225 degradation, k, was calculated from the first order plot (eq 3). 226

$$\ln\left(\frac{A_0}{A}\right) = kt \tag{3}$$

231

Where  $A_0$  is the initial absorbance, A is absorbance after a time 227 (t), and k is the first order rate constant. All photocatalytic 228 experiments were triplicated, and the rate constants were within 229 5% error limit. 230

### 3. Results and Discussion

**3.1. Characterization of Modified Precursor.** 3.1.1. 232 FT-IR Spectral Studies. The nature of reaction inter-233 mediates formed from acetic acid and EDTA modified 234 titania precursor was investigated using FT-IR spec-235 troscopy. Formation of an ionic Ti-EDTA complex 236 responsible for the low temperature crystallization of 237 heterojunctions was observed from this study. The dif-238 ference in frequency  $\Delta v$  between antisymmetric and 239 symmetric vibrations of C=O bonds are usually related 240 to the type of coordination (Figure 1) in metal car-241 F1 boxylates.<sup>26</sup> Acetic acid modified gel exhibits charac-242 teristic antisymmetric and symmetric C=O stretching 243 vibrations at 1532 and 1412 cm<sup>-1</sup>, respectively.<sup>27,28</sup> The 244  $\Delta \nu$  value of 120 cm<sup>-1</sup> observed is characteristic of a 245 bridged acetic acid titanium isopropoxide complex as 246 reported by Sanchez et al.<sup>28</sup> Antisymmetric and sym-247 metric C=O stretching vibrations of EDTA modified 248

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Figure 2. Differential scanning calorimetric pattern of titania gel. (a) Control sample, (b) 0.5 ED-TiO<sub>2</sub>, (c) 1.0 ED-TiO<sub>2</sub>, (d) 1.5 ED-TiO<sub>2</sub>, (e) 2.0 ED-TiO<sub>2</sub>.

gel were found at 1625 and 1392 cm<sup>-1</sup>, respectively. The  $\Delta \nu$  value of 233 cm<sup>-1</sup> observed for the EDTA modified gel confirmed the monodentate and ionic behavior of COO group in the complex.<sup>29</sup> The spectrum of this complex was consistent with that of [Ti (H<sub>2</sub>O) (edta)] as observed by Sato et al.<sup>30</sup>

All EDTA modified samples showed identical FT-IR 255 spectra. The formation of the acetic acid bridged complex 256 was not observed in compositions containing both acetic 257 acid and EDTA. These results indicate a preferential for-258 mation of the [Ti (H<sub>2</sub>O) (edta)] complex from a mixture of 259 acetic acid, EDTA, and titanium tetraisopropoxide, which 260 can be correlated to the stronger ligand character of EDTA. 261 Variations of ionic character among the different EDTA 262 modified compositions were also investigated. An increase 263 of ionic character was observed as a decrease in antisym-264 metric stretching frequency values of C=O group (Sup-265 porting Information 1). A minimum antisymmetric value 266 of 1600 cm<sup>-1</sup> observed for 1:1 EDTA complex indicates it 267 is of the highest ionic character, leading to a lowest ther-268 mal stability. This was later confirmed by the completion 269 of phase transformation to rutile at 700 °C (From XRD). 270 Bridged acetate groups in the acetic acid complex effec-271 tively alter the condensation pathway and promote the 272 formation of linear polymers. This leads to the formation 273 of anatase titania at higher temperatures, whereas EDTA 274 275 complex exists as monomers resulting in the low tempera-276 ture crystallization of anatase and rutile. These results thus confirmed the formation of a high temperature stable cova-277 lent acetic acid complex and a thermally less stable ionic 278 EDTA complex of  $Ti^{4+}$  (Supporting Information 2). 279

3.1.2. Thermal Stability Analysis of Modified Precursor. Both the amorphous to anatase transition and the
 anatase-rutile transformation (ART) from acetic acid and

EDTA derived complexes of titania were analyzed using 283 differential scanning calorimetry (DSC) and thermo-284 gravimetric analysis (TGA) techniques. For both sam-285 ples, evaporation of water and acetic acid molecules from 286 the gel was observed as an endothermic DSC peak at 287 50 °C (Figure 2). The acetic acid complex showed only 288 F2 one exothermic peak at 360 °C representing the crystal-289 lization of anatase phase. Anatase to rutile transforma-290 tion was not observed for this complex up to 600 °C. 291 Removal of coordinated water molecules from the EDTA 292 complexes were observed as an endothermic peak at 293 265 °C. Since there are no such coordinated water mole-294 cules in acetic acid complex, a corresponding peak was 295 not observed in these samples. Among the different 296 EDTA complexes, the most ionic (1:1 EDTA) material 297 shows the lowest anatase and rutile crystallization tem-298 peratures of 320 and 498 °C respectively. Slight increases 299 in both temperatures were observed for complexes con-300 taining excess EDTA (1:1.5 and 1:2). This may be due to 301 the hindering effect of excess EDTA toward anatase 302 phase evolution and its transformation to rutile. An 303 additional exotherm responsible for the decomposition 304 of unreacted EDTA was also observed at 290 °C in 305 complexes containing excess EDTA (1:1.5 and 1:2).<sup>31</sup> 306 The formation of anatase-rutile heterojunctions were 307 observed only from EDTA modified titania precursor. 308 In contrast, acetic acid modified precursor shows higher 309 thermal stability. It is thus clear that in the case of EDTA 310 modified samples, amorphous to crystalline as well as 311 anatase to rutile transformations happened at lower 312 temperatures. 313

The total weight loss calculated for the acetic acid 314 modified complex from the TGA curve was 17%. An increased weight loss of 58% and 85% was observed for 1:0.5 316 and 1:2.0 EDTA modified compositions respectively 317

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Figure 3. XRD of samples calcined at 600 °C. (a) Control TiO<sub>2</sub>, (b) 0.5 ED-TiO<sub>2</sub>, (c) 1.0 ED-TiO<sub>2</sub>, (d) 1.5 ED-TiO<sub>2</sub>, (e) 2.0 ED-TiO<sub>2</sub> (A = anatase:  $\mathbf{R} = \text{rutile}$ ).

(Supporting Information 3). The weight losses for the 318 acetic acid complex happen in the range 30-250 °C and 319 320 300-500 °C because of the evaporation of solvents and the decomposition of the complex into crystalline anatase 321 titania. On the other hand, the EDTA complex decom-322 poses in three steps (30-150, 250-450, 530-600 °C), a 323 process corresponding to evaporation of solvents, decom-324 position of EDTA-titania complex in to anatase titania, 325 and anatase-rutile transformation, respectively. On in-326 creasing the amount of EDTA, a significant increased 327 weight loss in the anatase formation step was observed, 328 which can be due to the decomposition of excess EDTA 329 present in these samples. The lowering of both the anatase 330 331 and rutile formation temperature was also observed (from the TGA curve) on increasing the amount of 332 EDTA. 333

3.2. Formation of N-Doped Heterojunctions. 3.2.1. 334 X-ray Diffraction Studies. X-ray diffraction patterns 335 recorded from samples calcined at different temperatures 336 confirmed the high temperature stability of acetic acid 337 modified samples and low temperature crystallization 338 of N-doped anatase-rutile heterojunctions from EDTA 339 modified samples. The control sample showed only 340 the peaks characteristic of anatase phase up to 600 °C 341 (Figure 3), whereas all EDTA modified samples exist as F3 342 anatase-rutile mixtures at a temperature as low as 500 °C. 343 The crystallization of anatase-rutile heterojunctions were 344 observed even at 400 °C for 1:1 and 1:1.5 EDTA modified 345 samples (Figure 4). Complete rutile formation for the 1:1 F4 346 EDTA modified sample (which was found to be the most 347 ionic from FT-IR studies) was observed at 700 °C (Sup-348 porting Information 4). Compared to other EDTA modi-349 fied samples, 1:2.0 modified EDTA sample shows slow 350 anatase rutile transformation kinetics and contains 4% 351 352 anatase even at 800 °C. All samples calcined at 900 °C exist as 100% rutile (Supporting Information 5). Bridged 353 acetate groups in AcOH-Ti complex effectively alter the 354 condensation pathway and promote the formation of 355 linear polymers. This leads to the formation of anatase tit-356 ania at higher temperatures, whereas EDTA-Ti complex 357



Figure 4. XRD of samples calcined at 400 °C. (a) Control TiO<sub>2</sub>, (b) 0.5  $ED-TiO_2$ , (c) 1.0  $ED-TiO_2$ , (d) 1.5  $ED-TiO_2$ , (e) 2.0  $ED-TiO_2$  (A = anatase;  $\mathbf{R} = \text{rutile}$ ).

exists as monomers and results in the low temperature 358 crystallization of both anatase and rutile phases having 359 large crystallite sizes. Slight changes ( $2\theta = 0.2^{\circ}$ ) in the 360 anatase 110 peak positions for EDTA modified titania 361 samples were observed even at 400 °C compared to the 362 control sample (Supporting Information 6), which results 363 from changes in the lattice parameters as a result of 364 nitrogen doping.<sup>32</sup> Thus XRD results revealed the low 365 temperature crystallization of anatase rutile heterojunc-366 tions from EDTA modified titania samples and successful 367 lattice incorporation of nitrogen. 368

3.2.2. Raman Studies. Raman spectroscopy, a more 369 surface sensitive technique was employed as an additional 370 tool to confirm phase composition and surface homo-371 geneity.<sup>33</sup> The Raman active modes for anatase  $(A_{1g}+$ 372  $2B_{1g}+3E_{g}$  at 147, 197, 396, 516, and 638 cm<sup>-1</sup>) and rutile 373  $(A_{1g}+B_{1g}+B_{2g}+E_g \text{ at } 144, 238, 447, \text{ and } 611 \text{ cm}^{-1})$  were used as fingerprints.<sup>34-36</sup> The samples 1:1 EDTA and 374 375 1:1.5 EDTA calcined at 400 °C has peaks characteristic of 376 anatase rutile mixtures, while other samples show peaks 377 of pure anatase phase (Supporting Information 7). Only 378 the acetic acid modified sample gives peaks of anatase at 379 600 °C; all EDTA modified samples were anatase-rutile 380 mixtures (Figure 5). All samples calcined at 900 °C shows 381 F5 peaks at 144, 611, 447, and 238 cm<sup>-1</sup>, which confirmed 382 the existence of 100% rutile phase (Supporting Informa-383 tion 8). Thus the results observed from X-ray diffraction 384 studies were further confirmed by Raman spectroscopy. 385

3.2.3. FT-IR Spectral Studies. Spectra of heterojunctions obtained were highly dependent on the processing temperature and nature of dopant species (Figure 6). Considerable shifting of Ti-O-Ti stretching frequencies to higher energy was observed for N-doped samples in comparison to the undoped one. At 400 °C, Ti-O-Ti stretching peak of N-doped sample was observed at a

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**Figure 5.** Raman spectra of samples calcined at 600 °C. (a) Control TiO<sub>2</sub>, (b) 0.5 ED-TiO<sub>2</sub>, (c) 1.0 ED-TiO<sub>2</sub>, (d) 1.5 ED-TiO<sub>2</sub>, (e) 2.0 ED-TiO<sub>2</sub>.



Figure 6. FT-IR Spectrum of calcined titania. (a) Control TiO<sub>2</sub> 400, (b) 0.5 ED-TiO<sub>2</sub> 400, (c) Control TiO<sub>2</sub> 600, (d) 0.5 ED-TiO<sub>2</sub> 600, (e) Control TiO<sub>2</sub> 700, (f) 0.5 ED-TiO<sub>2</sub> 700.

higher energy of 508 cm<sup>-1</sup> compared to 465 cm<sup>-1</sup> for un-393 doped one. Similar shift in Ti-O-Ti stretching energies 394 were also observed for samples calcined at higher tem-395 peratures. These types of shifts to higher energies were 396 previously reported and assigned to the formation of 397 O-Ti-N and N-Ti-N bonds in N-doped titania.<sup>37</sup> 398 Since we observed the formation of N-Ti-N bonds 399 as an additional peak at 415 cm<sup>-1</sup> in samples calcined at 400 700 °C,<sup>37</sup> shifting observed for Ti-O-Ti stretching en-401 ergies below this temperature can be assigned to the 402 403 formation of O-Ti-N bonds through partial replacement of oxygen. No evidence for interstitial nitrogen 404 doping and other nitrogen impurities were observed from 405 the spectra. The lattice incorporation of nitrogen in two 406 different forms was thus observed from FT-IR results. 407 Formations of O-Ti-N and N-Ti-N bonds were also 408 confirmed through XPS studies and will be discussed in 409 the following section. 410

J. Solid State Chem. 2006, 179, 1383–1393.

3.2.4. XPS Studies. XPS results revealed two types of 411 lattice doping (O-Ti-N and N-Ti-N) and a decrease of 412 nitrogen content with an increase of EDTA concentration 413 (Supporting Information 9). An increase of EDTA con-414 centration accelerates crystallite growth (from XRD re-415 sults), which can decrease the amount N-doping as a 416 result of surface and lattice strain relaxation. Decrease of 417 nitrogen content was also observed with an increase of 418 calcination temperature. This can be due to the crystallite 419 growth and escape of the dopant species formed through 420 the decomposition of Ti-EDTA complex at higher tem-421 peratures. The N 1s binding energy peak of N-doped TiO<sub>2</sub> 422 heterojunctions were found to be highly depends on the 423 calcination temperature. In the case of samples calcined 424 at 700 °C, formation of N-Ti-N bonds were observed as 425 a peak at 396 eV (which corresponds to an additional FT-426 IR peak at  $415 \text{ cm}^{-1}$ ) (Figure 7 b).<sup>38</sup> This can be due to the 427 F7 formation of specific Ti-N bonds through an oxygen 428 vacancy compensation mechanism at a higher tempera-429 ture. Only one peak was observed at 400.1 eV in samples 430 calcined at 600 °C (Figure 7 a), which is a value greater 431 than the typical N 1s binding energy of 397.2 eV in TiN.<sup>39</sup> 432 Since FT-IR and XRD results indicate lattice doping, this 433 peak therefore can be attributed to the 1s electron binding 434 energy of the N atom in the environment of O-Ti-N in 435 lattice N-doped titania. The binding energy shifting can 436 be understood by the fact that the N 1s electron binding 437 energy is higher when the formal charge of N is more 438 positive (e.g., 408 eV in NaNO<sub>3</sub>), compared to zero or a 439 negative formal charge (398.8 eV in NH<sub>3</sub>).<sup>39,40</sup> When a 440 nitrogen substitutes for the oxygen in the initial O-Ti-O 441 structure, the electron density around N is reduced com-442 pared to that in a TiN crystal (because of the O atom on 443 the Ti atom). Thus the N 1s binding energy in an 444 O-Ti-N environment (400.1 eV) is higher than that in 445 an N-Ti-N environment (396 eV) where the N atom 446 replaces the O atom. These observations are consistent 447 with the earlier XPS characterization results associated 448 with the oxidation of TiN surfaces by Saha and 449 Tomkins.39 450

As reported by Saha and Tomkins, pure titania sample 451 has a typical Ti 2p binding energy of 459.4 eV.<sup>39</sup> A dec-452 rease of 0.35 and 0.96 eV in Ti 2p binding energy was 453 observed for N-doped heterojunctions obtained at 600 454 and 700 °C, respectively, in comparison to the standard 455 sample (Figure 8). A decrease of Ti 2p binding energy is a 456 F8 direct measure of the lowering of the valence state level of 457  $Ti^{4+}$  to  $Ti^{3+}$  and  $Ti^{2+}$  level as a result of the lattice 458 substitution of nitrogen for oxygen.<sup>35,41</sup> Lattice incor-459 poration of nitrogen creates Ti-N bonds by the partial 460 replacement of  $O^{2-}$  with N<sup>-</sup>. This results in an electron 461 density decrease and partial reduction of Ti<sup>4+</sup> to Ti<sup>3+</sup> and 462  $Ti^{2+}$ , which reflects as a decrease in Ti 2p binding 463

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Figure 7. N 1s peaks in the XPS plot of calcined titania. (a) 0.5 ED-TiO<sub>2</sub> 600, (b) 0.5 ED-TiO<sub>2</sub> 700.



Figure 8. Ti 2p peaks in the XPS plot of calcined titania. (a) Control  $TiO_2$  600, (b) 0.5 ED- $TiO_2$  600, (c) 0.5 ED- $TiO_2$  700.

energy.<sup>41</sup> An additional O 1s peak appeared for N-doped 464 heterojunctions at 532 eV (Supporting Information 10), 465 which was first observed by Saha and Tomkins and most 466 recently characterized by Gyorgy et al. in their depth 467 profiling study on TiN surfaces. Gyorgy et al. assigned 468 this feature to the formation of oxidized TiN, which leads 469 to the Ti-O-N structure.42 Our study suggests that 470 the appearance of this peak is a consistent feature for 471 the nitrogen substitution in TiO2 and signifies the forma-472 tion of O-Ti-N and N-Ti-N structures. Moreover, 473 Rodriguez et al. measured the O 1s peak for -NO and 474 -NO<sub>2</sub> at 533.5 eV,<sup>43</sup> compared to the 532 eV value in our 475 study, which can exclude the novel properties related to 476 surface adsorbed nitrogen species. The presence of ad-477 ventitious elemental carbon was also identified in samples 478 479 (peak at 285 eV), (Supporting Information 11) which is an

Table 1. Band Gap Values of Heterojunctions Calcined at Different Temperatures

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composition	band gap (eV) 400 °C	band gap (eV) 600 °C	band gap (eV) 700 °C		
control TiO <sub>2</sub>	3.11	3.16	3.15		
$0.5 \text{ ED-TiO}_2$	3.03	3.02	2.98		
$1.0 \text{ ED-TiO}_2$	3.00	3.00	2.96		
$1.5 \text{ ED-TiO}_2$	2.98	2.98	2.95		
$2.0 \text{ ED-TiO}_2$	3.05	2.97	2.93		

unavoidable presence on all air-exposed materials.<sup>44</sup> It is therefore clear from the above observations that the lowering of Ti 2p binding energies and additional O 1s peak observed for N-doped heterojunctions together with FT-IR and XRD results well explained successful lattice incorporation of nitrogen through the formation of O-Ti-N and N-Ti-N bonds.

3.2.5. UV/vis Spectroscopic Studies. Band gap values 487 calculated through diffuse reflectance measurements also 488 support the variations of nitrogen and rutile concentra-489 tions in heterojunctions. All N-doped heterojunctions 490 show narrowing of the band gap in comparison to the 491 control sample (Table 1). The narrowing effect was com-492 T1 paratively high for the composition  $0.5 \text{ ED-TiO}_2$ , which 493 contain the highest amounts of nitrogen at each calcina-494 tion temperature (Figure 9). Band gap narrowing effect 495 F9 was very low for compositions containing higher amounts 496 of EDTA. This can be due to decreased N-doping as a 497 result of excessive crystallite growth on increasing EDTA 498 concentration. It was previously reported that the nitro-499 gen doping can lead to a mixing of N 2p orbitals with O 2p 500 orbitals to form intermediate energy levels and shift the 501 absorption edge toward visible light region.<sup>14,45</sup> Nitrogen 502 doping also accelerate the formation of oxygen vacancies, 503 which form additional energy levels below the conduction 504 band. This promotes electronic excitation from the 505 valence band to the additional energy level by absorbing 506 visible light.<sup>46</sup> A lowering of band gap values through 507

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Figure 9. Diffuse reflectance spectra of (a) Control  $TiO_2$  600, (b) 0.5 ED- $TiO_2$  600, and (c) 2.0 ED- $TiO_2$  600.

both of these mechanisms highly depends on the dopantconcentration, which explains the band gap variationsobserved above.

511 The amount of rutile also plays a significant role to shift the absorption edge of heterojunctions toward the 512 visible light region. Since it is evident from the XPS 513 results that the amount of nitrogen decreases on in-514 creasing the EDTA content, a decrease in the band gap 515 after the composition 0.5 ED-TiO<sub>2</sub> happens mainly 516 because of rutilation. For this reason, all samples show 517 a decrease in the band gap on increasing the calcination 518 temperature. At 400 °C, band gap narrowing effect was 519 low because of less rutilation at lower temperatures. The 520 superior visible light absorbing capability of hetero-521 522 junctions and their dependence on the amount of nitrogen and rutile were thus verified from diffuse reflectance 523 results. 524

3.2.6. Surface Area Measurements. Isotherms of hetero-525 junctions and the control sample show type IV char-526 acteristics with H1 type (cylindrical shape) of hysteresis 527 (Figure 10).<sup>47</sup> The high steepness hysteresis loop ending at F10 528 a relative pressure  $(P/P_0)$  of 0.6 is a measure of high order 529 of mesoporosity in these samples.<sup>48</sup> Degussa P-25, the 530 standard photocatalyst, has a type II isotherm with 531 small hysteresis behavior (Figure 11). This fact demon-F11 532 strates the presence of a large quantity of agglomerated 533 particles formed through a diffusion process, which 534 decreases the porosity. Among the different samples 535 compared, the control sample has the highest surface area 536  $(134 \text{ m}^2/\text{g})$  and pore volume (0.209 cc/g) at 400 °C (Table 2). T2 537 All heterojunction samples have lower surface area com-538 pared to the control sample at all calcination temperatures. 539 For both heterojunctions and the control sample, a con-540 siderable decrease in surface area and pore volume was 541 observed with an increase in calcination temperature. 542



Figure 10.  $N_2$  adsorption-desorption isotherm and the pore size distribution (inset) of 1.5 ED-TiO<sub>2</sub> 400.



Figure 11.  $N_2$  adsorption-desorption isotherm and the pore size distribution (inset) of Degussa P-25.

Table 2. Textural Properties of Heterojunctions Calcined at Different Temperatures

composition	surface area (m²/g)	pore diameter (nm)	pore volume (cc/g)
control 400	134.2	4 65	0.209
control 600	40.06	8.16	0.104
1.5 ED-TiO <sub>2</sub> 400	79.6	5.64	0.168
1.5 ED-TiO <sub>2</sub> 600	21.28	8.01	0.052
Degussa P-25	46.7	3.89	0.129

An increase in pore diameter was also observed with an 543 increase in calcination temperature, which confirms the 544 presence of interstitial pores and its grain growth.<sup>47</sup> 545 Among the samples calcined at 400 °C, the most active 546 catalyst (1.5 ED-TiO<sub>2</sub>) showed a larger pore diameter 547 (5.6 nm) than the control sample (4.6 nm) because of the 548 large grain size resulting from the thermally less stable 549 titania precursor. The same composition has a smaller pore 550 size at 600 °C compared to the control sample. This can be 551 due to the densification of powder resulting from excessive 552 grain growth. Even though the heterojunctions have poor 553

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**Figure 12.** Transmission electron micrograph of 1.5 ED-TiO<sub>2</sub> 400. (a) TEM, (b) HRTEM, and (c) SAED pattern.

textural properties in comparison to that of the con-554 trol sample, properties of the most active heterojunction 555  $(1.5 \text{ ED-TiO}_2 400)$  were superior to that of Degussa P-25. 556 Better textural properties of the control sample arise 557 because of the formation of thermally stable acetic acid-558 Ti<sup>4+</sup> complex,<sup>28</sup> and lower textural properties of hetero-559 junctions result from poor thermal stability of ionic 560 intermediate formed by the reaction between Ti<sup>4+</sup> and 561 562 EDTA.

3.2.7. Transmission Electron Microscopic Studies 563 (TEM). The nanocrystalline structure and phase compo-564 sition of the most active heterojunction  $(1.5 \text{ ED-TiO}_2)$ 565 400) had been investigated by combined high resolution 566 567 transmission electron microscopy (HRTEM)-selected area electron diffraction (SAED) analysis as shown in 568 Figure 12. The sample was found to be consisting of nano-F12 569 particles having two different sizes. The average particle 570 size of about 90% of the particles was  $10 \pm 3$  nm, and 571 the remaining 10% were about  $15 \pm 3$  nm in size. These 572 results were consistent with the average crystallite size 573 and anatase-rutile ratio obtained from the XRD analysis 574 (90% 13 nm sized anatase and 10% 18 nm rutile). The 575 15 nm sized rutile nanoparticles were found to be dis-576 tributed between anatase nanoparticles and thereby faci-577 litate the formation of heterojunctions (Figure 12 a, b). 578 The selected area electron diffraction (SAED) pattern 579 contains a series of Debye-Scherrer rings that corre-580 spond to an anatase-rutile mixture (Figure 12 c). The 581 d-spacing values 3.38, 2.23, 1.70, and 1.22 Å calculated 582 from the pattern represent the anatase lattice planes with 583 Miller indices (1 0 1), (0 0 4), (1 0 5), and (2 0 4), re-584 spectively.<sup>49,50</sup> The rutile lattice planes (1 1 0), (1 0 1), 585 (111), and (220) were also identified from the correspon-586 ding *d*-spacing values of 2.90, 2.5, 2.1, and 1.59 Å.<sup>49,50</sup> The 587 complex SAED ring pattern confirmed the presence of 588 polycrystalline anatase and rutile nanoparticles oriented 589 in a random manner.<sup>50</sup> 590

**3.3.** Photocatalytic Studies. 3.3.1. Visible Light Photocatalytic Degradation of MB over  $TiO_2$  Heterojunctions. All N-doped heterojunctions were more visible light active than the control sample or EDTA modified samples with a single phase toward the photocatalytic degradation of methylene blue. The most active heterojunction 1.5 ED-TiO<sub>2</sub> 400 containing 90% of 13 nm anatase and 597 10% of 18 nm rutile shows over 9 times more visible light 598 activity compared to Degussa P-25 (Figure 13 a, b). Rate 599 F13 constants obtained for the most active catalyst and 600 Degussa P-25 were 0.038 and 0.004 min<sup>-1</sup>, respectively 601 (Figure 14). Significantly higher visible light activity of 602 F14 nitrogen doped anatase rutile heterojunctions can be 603 explained on the basis of band gap narrowing and effi-604 cient charge separation at the anatase-rutile interface. 605 Results obtained from XRD, FT-IR, XPS, and UV/ 606 vis spectroscopy confirmed the formation of N-doped 607 anatase-rutile heterojunctions having low band gap values. 608 Additional energy levels created as a result of efficient 609 doping can utilize visible light for the generation of 610 electron hole pairs. Asahi et al. explained the visible light 611 sensitization of N-doped TiO<sub>2</sub> on the basis of Ti-N 612 bonding. Irie et al. also prepared N-doped TiO<sub>2</sub> by a 613 NH<sub>3</sub>-treatment method.<sup>38</sup> They observed the N 1s peak 614 at 396 and 400 eV in the XPS analysis and ascribed the 615 396 eV state to visible light sensitization. However, the 616 results of Diwald et al. rather indicated a negative con-617 tribution of Ti-N bonding to photocatalytic activity.<sup>51</sup> 618 So, the contribution of Ti-N bonding to the visible-light 619 sensitization of TiO<sub>2</sub> has not been completely under-620 stood. In our experiments, all nitrogen doped anatase 621 rutile mixtures at 700 °C with lower nitrogen content 622 (containing N-Ti-N bonds) show better (except 1:0.5 623 ED-TiO<sub>2</sub>) photocatalytic activity compared to samples at 624 600 °C (containing O-Ti-N bonds) (Supporting Infor-625 mation 12). This behavior can be correlated to the 626 formation of higher amounts of photocatalytic enhancing 627 Ti-N bonds at a higher temperature. The discrepancy 628 observed in the case of 1:0.5 EDTA composition may be 629 due to the presence of higher amounts of rutile at 700 °C. 630

In spite of its better textural properties, the control 631 sample was found to have very low visible light activity 632 compared to the most active sample, which has poor text-633 ural properties but high visible light activity. It is there-634 fore evident from the photocatalytic studies that the 635 textural properties of catalysts are not the only factor 636 determining the reaction kinetics. It can be concluded 637 that the composition of heterojunctions and band gap 638 values contribute more toward the photocatalytic activity 639 in comparison to the textural properties. 640

Crystallite sizes of anatase and rutile phases were 641 smaller for the most active heterojunction (1.5 ED-TiO<sub>2</sub> 642 400) than that in Degussa P-25 (Supporting Information 643 14). This facilitates the formation of more heterojunc-644 tions and thereby increases the activity. It is known that 645 the conduction band edge of rutile is approximately 0.2 646 eV lower than that of anatase. On this basis, the model 647 that the photo excited electrons (from the visible midgap 648 level) are effectively transferred from the conduction 649 band of anatase to that of rutile has been proposed 650 (Figure 16). When a small amount of rutile phase coexists with anatase phase, the interface between anatase and 652

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<sup>Tanemura, S.</sup> *Appl. Surf. Sci.* 2003, *212*, 255.
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Figure 13. Absorption spectra of visible light induced methylene blue degradation using (a) Degussa P-25 and (b) 1.5 ED-TiO<sub>2</sub> 400.



Figure 14. Kinetic study of (a) Degussa P-25 and (b) 1.5 ED-TiO<sub>2</sub> 400.

rutile promotes the electron transfer, and rutile works as a 653 reaction site for photo reduction. Photo oxidation can 654 take place either on anatase or on the rutile surface. The 655 heterojunctions obtained at 700 °C containing more Ti-656 N bonds showed a higher photocatalytic activity in com-657 parison to those obtained at 600 °C. The most active 658 heterojunction was the one obtained at 400 °C, and its 659 highest activity was found to be as a result of optimum 660 anatase-rutile composition, nanocrystalline nature, and 661 nitrogen doping. A decrease in photocatalytic activity 662 was observed for heterojunctions synthesized at higher 663 temperatures (Supporting Information 12). This can be 664 due to the formation of a higher amount of rutile phase 665 having higher electron hole recombination rate, and 666 decreases in the amount of more photoactive anatase 667 phase, nitrogen content, and surface area. Higher blue 668 shift of methylene blue absorption peaks during the 669 photocatalytic experiment of heterojunction samples in-670 dicates faster dye degradation through a N-demethylation 671 mechanism.52,53 The higher visible light photocatalytic 672



**Figure 15.** Photo luminescence spectra of samples (a)  $1.5 \text{ ED-TiO}_2 400$ , (b) Degussa P-25, (c) pure anatase, and (d) pure rutile.



Figure 16. Electron transfer mechanism in N-doped anatase rutile heterojunction.

activity observed for the heterojunctions can be therefore, 673 correlated to the combined band gap narrowing effect of 674 the nitrogen and electron hole separating effect of rutile. 675

*3.3.2. Luminescence Studies.* Information regarding 676 the presence of surface states, formation of photo induced 677 charge carriers, and their recombination kinetics can be drawn from the photoluminescence (PL) spectrum of a 679

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semiconductor material.<sup>54</sup> The phenomenon of photo-680 catalysis involves a competitive formation of electron 681 hole pairs and their recombination (radiative or non-682 radiative). Since defect concentration and life times of 683 photoexcited species play an important role in photoca-684 talysis, investigation of photocatalysts through PL spec-685 troscopy is important to obtain critical reasons behind the 686 enhanced photocatalytic activity. A strong correlation 687 between PL intensity and photocatalytic activity was 688 established by previous research.<sup>54,55</sup> 689

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Among different samples, pure rutile has highest band-690 band PL intensity followed by pure anatase, Degussa 691 P-25, and 1.5 ED-TiO<sub>2</sub> 400 (Figure 15). Band-band PL intensities mainly depend on the extent of photoinduced 693 charge carrier separation. The lower band gap and higher 694 surface defects of rutile sample accelerate charge carrier 695 recombination, which reflects as the most intense band-696 band PL peak.<sup>56</sup> Efficient electron hole separation result-697 ing from a wider band gap and smaller crystallite size 698 makes anatase less luminescent in comparison to rutile. 699 The lower PL intensity of Degussa P-25 (which is a 700 mixture of 70% anatase and 30% rutile) can be explained 701 on the basis of better electron hole separation resulting 702 from the electron transfer from the conduction band of 703 anatase to that of rutile.7 The slight red shift of the 704 band-band transition observed for Degussa P-25 may 705 be due to the presence of more rutile. Among the different 706 N-doped heterojunctions, the lowest PL intensity was 707 observed for sample 1.5 ED-TiO<sub>2</sub> 400. The optimum 708 phase composition (90% anatase +10% rutile) and nano-709 crystalline nature of components (which facilitates the 710 formation of more heterojunctions) helps the effective 711 712 transfer of photo excited electrons from anatase to rutile and slows down the recombination process. Also photo-713 excited electrons formed on rutile can be transferred to 714 the defect states of anatase, promoting effective charge 715 carrier separation. These processes are responsible for the 716 lower PL intensity and very high photo activity of EDTA 717 modified sample (1.5 ED-TiO<sub>2</sub> 400). Excitonic intensities 718 also follow the same order of band-band intensities 719 because of the same reasons. No additional PL bands 720 from nitrogen impurities were observed for heterojunc-721 tions, which eliminated the presence of surface adsorbed 722 723 nitrogen impurities. 724

Lower band-band and excitonic PL intensities resulting from efficient charge separation were reported for 725 TiO<sub>2</sub> and ZnO doped with Sn, W, Ag, and Au.<sup>55,57-61</sup> 726 These semiconductors showing lower PL intensities were 727 also reported to exhibit higher photocatalytic activity.<sup>59,62</sup> 728 In short, any electron capturing agents can lower both the 729

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band-band and the excitonic PL intensities, which results 730 in an increase in photocatalytic activity. Thus, the lowest PL 731 intensity and the very high visible light activity of nitrogen 732 doped anatase rutile heterojunctions can be explained on 733 the basis of efficient charge separation at the anatase-rutile 734 interface. It is known that the conduction band edge of rutile 735 is approximately 0.2 eV lower than that of anatase.<sup>7,18</sup> On 736 this basis, the model that the photoexcited electrons (from 737 the visible midgap level) are effectively transferred from the 738 conduction band of anatase to that of rutile has been 739 proposed. 740

### 4. Conclusions

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Nitrogen doped anatase-rutile heterojunctions having 742 high visible light photocatalytic activity were developed 743 through an ionic intermediate [Ti (H<sub>2</sub>O) (edta)]. Low 744 temperature crystallizations of heterojunctions were 745 characterized using DSC, TGA, XRD, HRTEM, and 746 Raman techniques. The nature of dopants was investi-747 gated using FT-IR spectroscopy, XRD, and XPS techni-748 ques. Conclusive evidence for the formations of both 749 O-Ti-N and N-Ti-N bonds were observed from FT-750 IR and XPS studies, and substitutional nitrogen doping 751 was found to have an enhancing effect on the photocata-752 lytic activities of heterojunctions. The most active 753 N-doped heterojunction obtained at 400 °C showed a 754 9-fold enhancement for the visible light decomposition of 755 methylene blue in comparison to the commercial catalyst 756 Degussa P-25. The heterojunction has lower lumines-757 cence intensity as a result of better electron hole separa-758 tion, and better textural properties in comparison to 759 Degussa P-25. The highest photocatalytic activity of the 760 optimum sample (1.5 ED-TiO<sub>2</sub> 400) was due to the com-761 bined effect of better electron hole separation, a lower 762 band gap resulting from effective nitrogen doping, and 763 nanocrystalline nature compared to the standard photo-764 catalyst. 765

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Supporting Information Available: Further details are given in 774 the supporting tables 1-3 and figures 1-13. This material is 775 available free of charge via the Internet at http://pubs.acs.org. 776

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