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Journal:	<i>Faraday Discussions</i>
Manuscript ID	FD-ART-11-2018-000191.R1
Article Type:	Paper
Date Submitted by the Author:	04-Feb-2019
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Cite this: DOI: 10.1039/xxxxxxxxxx

# Influence of Carbonaceous Species on Aqueous Photo-catalytic Nitrogen Fixation by Titania<sup>†</sup>

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Accepted Date

DOI: 10.1039/xxxxxxxxxx

www.rsc.org/journalname

For decades, reports have suggested that photo-catalytic nitrogen fixation by titania in an aqueous environment is possible. Yet consensus does not exist regarding how the reaction proceeds. Furthermore, the presence of an aqueous protonated solvent and the similarity between the redox potential for nitrogen and proton reduction suggest that ammonia production is unlikely. Here, we re-investigate photo-catalytic nitrogen fixation by titania in an aqueous environment through a series of photo-catalytic and electrocatalytic experiments. Photo-catalytic testing reveals that mineral phase and metal dopants play a marginal role in promoting nitrogen photofixation, with ammonia production increasing when the majority phase is rutile and with iron dopants. However, the presence of trace amount of adsorbed carbonaceous species increased the rate of ammonia production by two times that observed without adsorbed carbon based species. This suggests that carbon species play a potential larger role in mediating the nitrogen fixation process over mineral phase and metal dopants. We also demonstrate an experimental approach aimed to detect low-level ammonia production from photo-catalyst using rotating ring disk electrode experiments conducted with and without illumination. Consistent with the photocatalysis, ammonia is only discernible at the ring with the rutile phase titania, but not with the mix-phased titania. Rotating ring disk electrode experiments may also provide a new avenue to attain a higher degree of precision in detecting ammonia at low levels.

## 1 Introduction

Reports of photocatalytic nitrogen fixation on titania have surfaced for nearly eight decades<sup>1</sup>. Demonstrations have occurred in both the gas and aqueous phase, and with various environmental conditions (relative humidity, temperature, pressure). Nearly all experimental observations suggest that abiotic nitrogen photofixation may be possible at ambient temperature and pressure<sup>2,3</sup>. This is impactful as it promotes the possibility for nutrient based fertilizer production from environmentally abundant materials (minerals) using only the sun as a source of energy. Considering the state-of-the-art thermochemical approach for fertilizer production is responsible for consuming 1% of global energy and emitting 3% of global CO<sub>2</sub>, photocatalytic nitrogen fixation could have a significant influence on improving the environmental impact associated with ammonia production<sup>2</sup>.

Photocatalytic nitrogen fixation also has environmental significance, as nitrogen fixation is a critical catalytic process in the nitrogen cycle. Although, biological fixation is the most notable natural entry point for dinitrogen in soils, additional entry points driven through photon based reactions on earth abundant minerals are not outside the realm of possibility. A prominent Indian soil scientist (N. Dhar) in the 1940s suggested that the generation of fixed nitrogen in soils was possible through light based interactions<sup>1</sup>. Schrauzer and Guth explored this hypothesis further in the late 1970s through conducting a series of experiments aimed at discerning the photocatalytic nitrogen fixation activity on environmental catalyst (global sands)<sup>3-5</sup>. This work concluded that the presence of titania in sands was responsible for nitrogen photofixation. Despite these intriguing findings, the potential impacts of photocatalytic nitrogen fixation on the nitrogen cycle is still limited to only back-of-the envelope based calculations<sup>6</sup>.

This slow progress is due to a lack of understanding regarding the catalytic processes that enable nitrogen fixation to occur on titania based photocatalyst. The wide band gap mineral coupled with the location of the band edges, suggest that nitrogen reduction is unlikely. Furthermore, in environments where hydrogen evolution is possible (aqueous conditions), selectivity toward

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<sup>†</sup> Electronic Supplementary Information (ESI) available: [details of any supplementary information available should be included here]. See DOI: 10.1039/b000000x/

nitrogen reduction would require a low overpotential. Furthermore, the low measurable yields, high propensity for contamination and numerous sources of measurement error have all contributed to the significant debate regarding the probability of photocatalytic nitrogen fixation occurring on titanium dioxide<sup>7–11</sup>. Most prior studies have emphasized the role iron dopants and oxygen vacancies play in activating dinitrogen on titania<sup>12–14</sup>. Significant evidence supports the hypothesis that iron dopants aid in stabilizing oxygen vacancies, and that the presence of vacancies weakens the dinitrogen triple bond, enabling dinitrogen reduction. While possible, a more recent hypothesis suggests that carbon may play a critical role in nitrogen photofixation on titania<sup>15</sup>. Specifically, gas phase AP-XPS based experiments demonstrated the presence of adventitious carbon was important for promoting interactions between titanium dioxide and nitrogen. The researchers hypothesized that a photocatalyzed carbon radical in concert with defects may promote nitrogen photofixation to ammonia. These gas phase experiments promote the need to understand if photocatalyzed carbon radicals also play a similar role in an aqueous photocatalytic environment.

Here, we aim to expand upon work completed in the gas phase; to explore the potential role carbonaceous species may play in promoting photo-catalytic nitrogen fixation in aqueous based experiments. While adventitious carbon and carbon dioxide may be the dominant carbonaceous species present in gas phase photocatalysis, here organic hole scavengers (methanol, ethanol, formic acid) are the most prominent sources of carbon. We begin by probing the photocatalytic activity of rutile, mixed phase and iron-doped titania in the presence and absence of carbonaceous species. Rotating ring disk electrode experiments conducted on the catalyst also aim to determine if ammonia is detectable under illumination. Finally, the surface based active sites (oxygen vacancies and lattice iron) are mapped using bulk and surface based characterization.

## 2 Materials and Methods

### 2.1 Catalysts preparation

A commercial rutile titanium dioxide and mixed phase titanium dioxide were purchased by US Research Nanomaterials Inc and Alfar Aesar (Haverhill, MA). Iron-doped titanium dioxide was prepared by standard sol-gel methods<sup>16</sup>. Titanium dioxide sols were prepared through the addition of 5 mL of titanium tetraisopropoxide (TTIP) solution into 50 mL of absolute ethanol. Then, the TTIP/ethanol solution added (dropwise) into 50 mL of distilled water which was adjusted to pH 1.5 with chloric acid under vigorous stirring at room temperature. After continuously stirring for 24 hrs, the resulting transparent solution was evaporated using a rotary heater at 50°C and dried in the oven at 70°C overnight. Resulting particles were calcined at 500°C for 1 hr under air, and stored in air tight vessels. Catalyst cleaning prior to use occur through a calcination process at 400°C for 4 hrs. This cleaning procedure prevented the detection of any ammonia during controls, which were conducted in the absence of nitrogen (argon gas and light) and in the absence of light (nitrogen gas and dark).

### 2.2 Photocatalysis Experiments

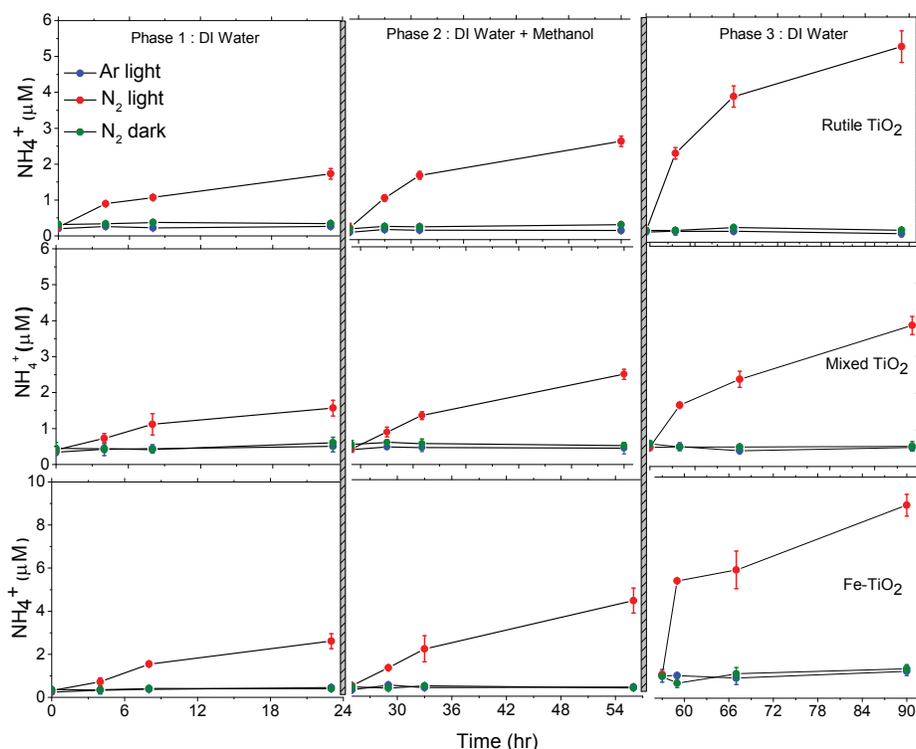
The photocatalytic nitrogen fixation experiments were conducted in a 500 mL reaction vessel at ambient temperature and atmospheric pressure. The temperature was fixed at 28°C during the tests and was monitored to ensure constant temperature existed under illumination. A 300 W Xenon-Mercury lamp (Newport Corporation - Irvine, CA) was used which contained a ultraviolet (UV) cutoff filter ( $\lambda > 320$  nm) and an infrared (IR) cutoff filter. In all tests, 300 mg of photocatalyst was suspended in 300 mL of aqueous electrolyte. The aqueous electrolyte consisted of deionized water, or a mixture of deionized water with a hole scavenger. Hole scavenger concentration remained fixed at five volume percent. Methanol, ethanol and formic acid were chosen as representative hole scavengers. The photocatalytic reactor was deaerated using nitrogen gas (purity $\geq$ 99.999%) which was bubbled through this solution for thirty minutes in the dark. Various times were investigated to ensure that the solution was completely saturated with nitrogen and that all residual dissolved oxygen was removed. After saturation was reached, the reactor was irradiated with ultra-violet light ( $\lambda > 320$  nm) from the top to the reactor. The reactor was continuously fed nitrogen gas during the entire twenty-four hour test period. Controls were conducted to probe the presence of contamination. In the first control, nitrogen gas was replaced by argon gas (purity $\geq$ 99.999%). All other conditions remained constant, including the degree of illumination. A second control was conducted in the dark with nitrogen gas. During all tests (including controls) approximately, 5 mL of the suspension was removed from the reaction vessel at given intervals to test for ammonium.

### 2.3 Ammonium measurement

Two milliliters of dilute acid (0.005 M dilute H<sub>2</sub>SO<sub>4</sub>) was added to the catalyst suspension removed from the photocatalytic reactor. The dilute acid was used to promote ammonia desorption from the catalyst surface<sup>4</sup>. The particle suspension were then filtered by 0.22 $\mu$ m precision-glide syringe needle (Sigma-Aldrich, St. Louis, Mo) and syringe filters (Hach Co). The desorbed ammonium concentration was analyzed by ionic chromatography using a cation exchange column (Aquion Dionex, Thermo Fisher). Ammonia was easily distinguished from other cations and had a retention time of 5.24 minutes. Standard solutions of ammonium chloride were used to create a calibration curve for the ion chromatograph. While many standard methods rely on optical measurements (UV-Vis) whereby an organic compound reacts with ammonia (indophenol blue), we found these methods were susceptible to erroneous data, and a lack of reproducibility, consistent with prior works<sup>17</sup>.

### 2.4 Electrochemical Characterization

Electrochemical testing was conducted using a rotating ring disk electrode experimental set up (Pine Research, Durham, NC). The catalyst was dispersed onto the disk, and the ring was poised to sense product (ammonia) formation. The catalyst was dispersed onto the disk using standard methods. A stock solution of 20% isopropanol and 0.5% Nafion solution is prepared by mixing 20



**Fig. 1** Ammonia yield with time during photo-catalytic testing under 300 W lamp. Experiments were conducted in three phases. Phase 1, 2 and 3 are displayed in panels 1-3. (a) Rutile titania (b) mixed phase titania and (c) iron-doped titania.

mL of isopropanol with 70 mL of distilled water and 10 mL of 5 wt% Nafion solution in a 100 mL volumetric flask. Then 25 mg of catalysts was added into the nafion/IPA solution and sonicated for one hour. After the suspension was well dispersed without aggregation or sedimentation, 10  $\mu$ L of the ink was drop cast onto the glassy carbon disk (6.2 mm diameter) and rotate in 1000 rpm for one hour using the air dry method<sup>18,19</sup>. A lamp (Asahi Spectra, Torrance, CA), was situated below the disk outside of the glass reactor. The lamp was used to irradiate the disk during experiments. A platinum ring (inner and outer diameter, 7 and 8.4 mm) served as the working electrode. A silver-silver chloride reference electrode (saturated potassium chloride) and a platinum wire counter electrode were also used. Cyclic voltammetry was performed while rotating the shaft of the working electrode at 500 rpm with a scan rate of 5 mV s<sup>-1</sup>. Simultaneously, the ammonia oxidation reaction was investigated at the ring. In all testing the supporting electrolyte consisted of sodium hydroxide. Alkaline conditions were deemed necessary in order to promote the ammonia oxidation reaction.

## 2.5 Material Surface Characterization

The morphology of the titanium dioxide materials were investigated by scanning electron microscopy (SEM) coupled with electron dispersive spectrometer (EDS). EDS coupled with XPS analysis confirmed that the iron-doped samples exhibited 0.69 atomic percent iron in the titania lattice. The as-prepared materials were identified through X-ray powder diffraction (XRD) using Penanalytical Xpert Pro Alpha-1 XRD system for crystalline identification

with Cu K $\alpha$  radiation ( $\lambda = 1.54051$ ). Step scanning was used with 2 $\Omega$  intervals from 10 $^\circ$  to 90 $^\circ$  with a residence time of 1 s. The chemical states of the surface elements were recorded using a x-ray photon spectrometer (XPS) by a Thermo K-Alpa XPS spectrometer equipped with monochromatic Al K $\alpha$  radiation as the X-ray source. The Ti2p, O1s, C1s, and Fe2p peaks were analyzed to access the surface properties (oxygen vacancies and dopants) of each catalyst.

## 3 Results and Discussion

### 3.1 Photocatalytic testing with carbonaceous species

Photocatalysis in an aqueous phase reactor was conducted with the three catalysts. Two off the shelf titanium dioxide catalyst were chosen. The first off-the-shelf catalyst had a mineral composition that was primarily rutile phase. The second off-the-shelf catalyst had a mineral composition that was a mixture of anatase and rutile phase. A third catalyst was synthesized in the lab and contained iron dopants at 0.69 atomic percent, as confirmed through energy dispersive x-ray spectroscopy and XPS. Iron was chosen as the representative metal dopant, due to the demonstrated high activity reported in prior photocatalytic nitrogen fixation investigations<sup>4,14</sup>.

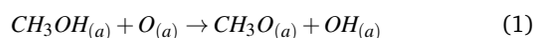
Photocatalytic experiments were conducted in three phases. In phase one, suspended particles were deaerated with either argon or nitrogen gas prior to illumination to remove all dissolved oxygen. The reactor was then illuminated for twenty-four hours, with samples removed at timed intervals. For all tests with argon, no ammonia was detected over the entire twenty four hour

test. Dark controls in the presence of nitrogen also yielded no detectable ammonia. This confirmed that little nitrogen based contamination was present on the catalyst or photo-catalytic reactor prior to and during testing. Thus all ammonia detected was a result of a light-based reaction.

During the phase one experiments, the rate of ammonia production did not change regardless of the catalyst (rutile, mixed phase, and iron-doped titanium dioxide). The rate of ammonia production ranged from  $0.25\text{--}0.38 \pm 0.03 \mu\text{M hr}^{-1} \text{g}^{-1}$ , indicating that the phase (rutile or anatase) and metal dopants (iron) did not play a significant role in improving the rate of nitrogen photofixation (Figure 1 - Phase 1). Prior investigations have highlighted that the phase (rutile)<sup>4</sup> and concentration of oxygen vacancies are critical material properties needed in order to increase the rate of ammonia production<sup>13</sup>. Furthermore, rutile phase catalyst have been shown both experimentally and theoretically to be active for both photo and electrocatalytic nitrogen fixation<sup>4,12</sup>.

Oxygen vacancies or point defects in the titanium dioxide lattice are also well known to exist in non-stoichiometric samples<sup>20</sup>. However, in the presence of oxygen or water, as is the case with aqueous-based photocatalytic experiments, vacancies can heal resulting in a near stoichiometric crystal. Thus, while oxygen vacancies may be an active site, the lifetime and concentration of this active site is uncertain in a aqueous environment. We also note that the initial rates observed are in line with prior investigations, which contained catalyst with a low concentration of oxygen vacancies<sup>13</sup>.

After the initial phase, the minerals were separated and recycled back to the reactor for phase two. In phase two, a hole scavenger was introduced into the system. Traditional hole scavengers are organic compounds (e.g. EDTA, methanol, ethanol) which can be easily oxidized at the valance band by photogenerated holes (Figure 1- Phase 2). Here, the aim in introducing a hole scavenger was not to minimize the rate of charge carrier recombination, but rather was to probe the hypothesis that nitrogen photofixation occurs due to an interaction with a photogenerated carbon radical. Methanol was chosen as a representative hole scavenger, as methanol is well studied with titania based photocatalytic systems, has been a hole scavenger used in prior nitrogen fixation investigations, and methanol has a well-documented reaction pathway<sup>21</sup>. Methanol oxidation is suspected to occur predominately through a dissociative pathway resulting in adsorbed methoxy species and hydroxyl groups<sup>22,23</sup>.



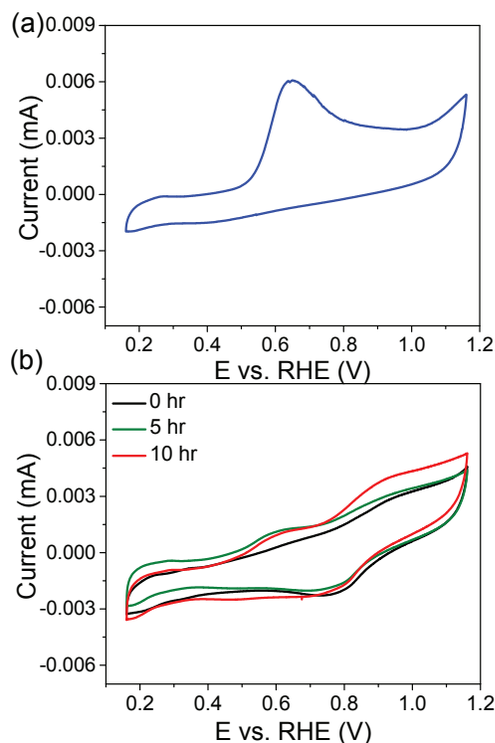
The methoxy species have furthermore been shown to generate carbon-based radicals that have been detected through both infrared and electronic paramagnetic resonance based spectroscopy<sup>22</sup>. With this goal in mind, phase two proceeded with methanol present in the deionized water. Methanol was supplied to the deionized water at 5 vol%, which is consistent with the concentrations traditionally supplied to photocatalytic systems, whereby methanol acts as a hole scavengers. We will also note that in addition to the carbon radicals generation due to the presence of methanol, a degree of adventitious carbon is also well

known to exist on all titania based photocatalyst.

For each catalyst, the rate of ammonia production increased on average by 60% when compared to testing conducted without a organic hole scavenger. With the rates increasing to  $0.4\text{--}0.65 \pm 0.04 \mu\text{M hr}^{-1} \text{g}^{-1}$ . This increase in the rate of ammonia produced is traditionally ascribed to the system no longer being limited by the hole driven reaction (water splitting). Instead of oxidizing water, methanol is oxidized by the large oxidative potential of the holes generated in titanium dioxide, preventing charge carrier recombination. In phase two, the performance of rutile, mixed phase and iron-doped titania was similar, indicating that phase and metal dopants again did not alter performance significantly. In addition, controls (nitrogen dark and argon light) resulted in no appreciable ammonia production.

In an effort to probe the role of the carbon radical more directly, the catalysts was recycled a third time back in deionized water which contained no methanol. Here, by recycling the catalyst exposed to methanol oxidation, the aim was to remove the methanol, while maintaining a degree of adsorbed methoxy groups (or other oxidized carbon species) on the catalyst. Prior investigations have shown that methanol desorbes easily from the catalyst and solution at 308-373 K, while the methoxy groups remain adsorbed ( $T_{\text{des}}=460 \text{ K}$ )<sup>23,24</sup>. Thus we hypothesize that in phase three, carbon radical generation may be possible in the absence of methanol due to these adsorbed species. In phase three, each catalyst experienced an increase in the rate of ammonia production (Figure 1- Phase 3) over testing with methanol (phase two), and more importantly over results observed with just deionized water (phase one). In addition, the rate of ammonia production differed between the three individual catalyst. The rate of ammonia produced on the iron-doped titania, rutile titania and mixed phase titania were  $1.3 \pm 0.07 \mu\text{M hr}^{-1} \text{g}^{-1}$ ,  $0.76 \pm 0.06 \mu\text{M hr}^{-1} \text{g}^{-1}$  and  $0.56 \pm 0.03 \mu\text{M hr}^{-1} \text{g}^{-1}$ . Thus the order of photocatalytic activity for the three catalyst was iron-doped > rutile > mixed phase (Figure 1). We also conducted control experiments (Rutile Titania) whereby catalyst were recycled three times without an organic scavenger. In these tests, that the ammonia yield remained at  $0.2 \pm 0.008 \mu\text{M hr}^{-1} \text{g}^{-1}$  (consistent with results obtained from Phase 1), indicating that the increased rate are not a result of catalyst recycling and handling.

Phase three photocatalytic testing clearly emphasizes that the adsorbed carbon radicals most likely play a role in mediating the nitrogen fixation process on titania, as the performance increase can not be ascribed to the presence of the hole scavenger. Additional testing also conducted with ethanol and formic acid resulted in similar trends (Figure S13), with slight improvements observed with formic acid. Prior theoretical insight suggests that the carbon radicals have strong-binding free energy (-1.89 eV) for nitrogen, which ultimately can aid in adsorbing and reacting dinitrogen<sup>12</sup>. Thus, the observed photocatalytic activity demonstrated here in an aqueous environment aligns with the prior gas phase experiments. With a range of reported values observed with photocatalytic nitrogen fixation based experiments on titania based photocatalyst<sup>4,7,13</sup>, here we highlight that in addition to material properties (bulk and surface), that carbon species prominently used within photocatalytic reactor may also

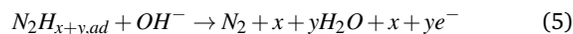
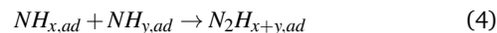
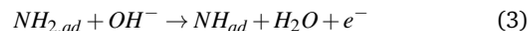
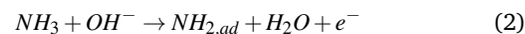


**Fig. 2** (a) Ammonia oxidation peak observed on the Pt ring with 0.1  $\text{NH}_4\text{Cl}$  in the supporting electrolyte. (b) Ammonia oxidation of rutile  $\text{TiO}_2$  under excitation of light and nitrogen gas.

influence the nitrogen fixation reaction mechanism and catalytic activity.

### 3.2 Electrocatalytic characterization of Titania

In addition to monitoring the ammonia production through bulk phase photocatalytic testing, we also probed the formation of ammonia through a series of rotating ring disk electrode (RRDE) experiments. The three catalyst (rutile, mixed phase and iron-doped titanium dioxide) were investigated through dispersing the catalyst on the disk, while the ring was then poised at the ammonia oxidation potential. The aim of the ring is to act as a sensor to detect products formed as a result of the photocatalyzed reactions (hole and electron driven) on the titanium dioxide. The disk therefore was alternatively excited and not excited by light that were situated below the catalyst. Experimental conditions were carried out with nitrogen and argon saturated electrolytes in the dark and light. A key difference between the photocatalytic experiments and the electrocatalytic experiments is the choice of electrolyte. While ideally, DI water would have been investigated, there were two chief challenges. The first being that ammonia oxidation in a highly resistive cell would produce very little current. With the goal to correlate photocatalyzed ammonia production to ring current, we needed to limit resistance through the use of a supporting electrolyte. Another challenge was that ammonia oxidation occurs through the following proposed mechanism on Pt:

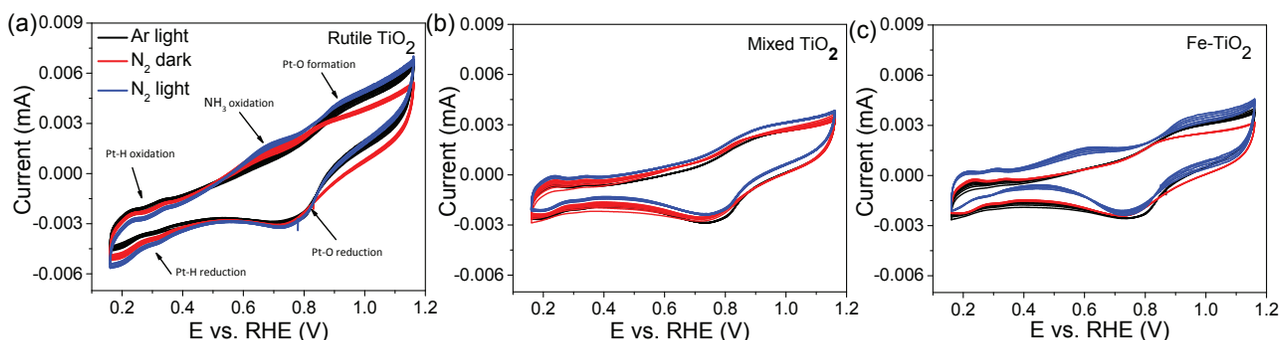


Since the ring utilized here was Pt, in order to promote the ammonia oxidation reaction,  $\text{OH}^-$  is needed in the supporting electrolyte at significant concentrations. For this reason an alkaline electrolyte was chosen, as prior investigations have shown that ammonia oxidation on Pt can not be detected in electrolytes with a pH lower than eight, and it is more preferable to operate with electrolytes that have a pH greater than ten<sup>25</sup>. It should be noted that titanium dioxide was not operated as an electrocatalyst, as titania was not electrochemically biased during the experiments.

In the control experiments, 0.1 M  $\text{NH}_4\text{Cl}$  was added to the supporting electrolyte and the ring voltage was swept from 0.2 vs. RHE to 1.2 vs. RHE. A noticeable peak was observed in all tests at  $\approx 0.7$  vs. RHE. This peak was attributed to ammonia oxidation and was used as a reference for future testing conducted with titanium dioxide (Figure 2a)<sup>26</sup>. Initial testing conducted with rutile titanium dioxide resulted in no peak during initial tests without light (Figure 2b - 0 hr). However, with the addition of light, the ammonia oxidation peak was observed. The catalyst was tested over a 10 hour period while illuminated, and the peak position did not vary significantly, while the peak current increased slightly. The extended experiment suggests that the rate of ammonia produced and oxidized remained constant with time (Figure 2b - 5, 10 hr). The other peaks besides ammonia oxidation are Pt-O formation (potential region 0.8 - 1 V), Pt-O reduction (potential region 1.2 - 0.7 V vs. RHE), Pt-H reduction (ca. 0.3 V vs. RHE) and Pt-H oxidation (ca. 0.35 V vs. RHE).

The three catalysts were first evaluated under similar conditions tested in the photocatalytic reactor. Namely, in an electrolyte saturated with nitrogen and argon gas, and under dark and light conditions. With the rutile phase titanium dioxide photocatalyst, the ammonia oxidation peak (ca. 0.7 V vs. RHE) was observed with nitrogen gas and the excitation of light (Figure 3a). No ammonia oxidation peak was present when nitrogen was replaced by argon, or when the sample was not illuminated (dark testing with nitrogen gas). In the potential region from 0.8 - 1.0 V vs. RHE, a Pt-O peak was observed. This is due to the presence of oxygen being formed under oxidative potentials on Pt. Furthermore, when scanning negatively in the potential region from 1.2 - 0.7 V vs. RHE, Pt-O reduction occurred. Pt-H desorption peak was showed in the potential region around 0.3 V vs. RHE under the negative scan and its redox couple was present around 0.35 V vs. RHE due to the Pt-H oxidation<sup>18</sup>. These peaks were consistent with all catalyst, which is to be expected as the reactions at the Pt based reactions are independent of the disk catalyst.

When the mixed phase titanium dioxide was used as the catalyst, no ammonia oxidation peak was visible under nitrogen (light and dark) and argon (light). Prior investigations have suggested that a mixed phase photocatalyst is non-ideal for nitrogen fixation, as rutile may be the most active phase. Furthermore, mixed



**Fig. 3** Rotating ring disk electrode experiments with and without excitation of light in argon and nitrogen atmospheres with (a) rutile (b) mixed phase and (c) fe-doped titania

phase photocatalyst are most active for hydrogen evolution. Since hydrogen evolution competes with nitrogen reduction it is not surprising that ammonia was not detected with the mixed phased photocatalyst (Figure 3b). With the iron-doped titania sample a more obvious ammonia oxidation peak was observed under nitrogen in the light. Again, controls with argon and nitrogen (dark) did not result in a peak (Figure 3c). The observed peak current (ammonia oxidation peak) measured in the ring electrode were in-line with the photocatalytic testing activity, which suggests that rutile phase and iron-doped titania are active for nitrogen fixation, whereas the mixed phase photocatalyst is the least active catalyst. The current and the charge from the ammonia oxidation area can be used to estimate the rate or nitrogen photofixation. The observed rates for the rutile and iron-doped titanium dioxide were on the order of  $10^{-6}$  mol per second, whereas the mixed phase catalyst was on the order of  $10^{-8}$  mol per second. Furthermore, the cyclic voltammetry of Pt disc electrode under nitrogen and argon gases flow, with and without excitation of light are shown in Supporting Information Figure S11. As shown, the Pt-O formation/reduction and Pt-H oxidation/reduction remained, but no ammonia oxidation are found with these three conditions indicating that Pt does not participate in ammonia formation with light illumination. Also, the most active iron-doped catalyst has been selected to perform the same phases 1-3 used during photocatalytic testing. The results were shown in Figure SI2. In phase one, iron-doped  $\text{TiO}_2$  was examined under nitrogen flow and excitation of light. After phase one, the disk is recycled back in electrolyte which contains methanol. In phase two methanol is oxidized when the disk is illumination (presumably generating carbon radicals). The methanol oxidation peak was observed at 0.8 V vs. RHE. In Phase 3, the electrolyte is replaced (no methanol), while adsorbed-methoxy-group remain. The result showed that the residual adsorbed surface groups again increased the ammonia formation, observable through the increase in the peak area (ca. 0.7 V vs. RHE). The observed rate of ammonia oxidation in phase three is 2 times higher than phase one. Also, the CV showed that no remain adsorbed methanol on the surface of catalyst since there is no methanol oxidation presented in the phase three experiment.

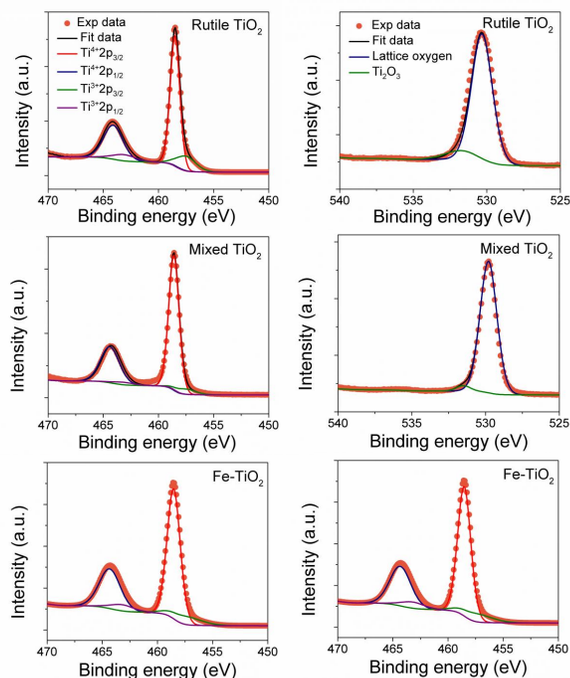
### 3.3 Catalyst Surface and Bulk Characterization

Both the photocatalytic testing and rotating ring disk electrode experiments provide evidence that the rate of nitrogen photofixation is greatest on the iron-doped titanium dioxide, followed by the rutile phase catalyst and the mixed phase catalyst. Material characterization was obtained by x-ray photoelectron spectroscopy (XPS) and X-ray powder diffraction (XRD). XPS provided insight into the degree of oxygen vacancies that were present in the catalyst, and the degree of iron dopant at the surface of the catalyst. It should be noted that the ex-situ analysis provides indirect evidence regarding the likelihood that a given catalyst has oxygen vacancies. It is not capable of determining how stable the vacancies are when immersed in the aqueous solution.

For the XPS, the Ti2p, O1s, C1s, and Fe2p regions were all analyzed. Determination of the degree of oxygen vacancies can be observed through monitoring the differences in the O1s and Ti2p regions (Figure 4). The O1s spectra typically contains two sub-peaks, one which represents the lattice oxygen and a second which details the surface  $\text{Ti}_2\text{O}_3$  groups. The Ti2p is typically fit with two peaks  $\text{Ti}2p_{3/2}$  and  $\text{Ti}2p_{1/2}$  (Figure 4b). In a well coordinated sample with no oxygen vacancies these peaks are sharp and are assigned to  $\text{Ti}^{4+}$  atoms. As the  $\text{Ti}2p_{3/2}$  and  $\text{Ti}2p_{1/2}$  broaden, two additional peaks can be introduced which are assigned to  $\text{Ti}^{3+}$  sites. Thus the two signatures of oxygen vacancies is the increase in  $\text{Ti}_2\text{O}_3$  groups (Figure 4a) and increase in the concentration  $\text{Ti}^{3+}$  sites sites. For the photocatalyst tested, we see that the most active samples have a higher degree of these two signatures, indicating that in the dry state, the amount of oxygen vacancies are indeed higher<sup>27,28</sup>.

According to the XPS analysis of Ti2p, we observed that the area of  $\text{Ti}^{4+}$  is 80% and  $\text{Ti}^{3+}$  is 20% in rutile  $\text{TiO}_2$ . Meanwhile, the area of  $\text{Ti}^{4+}$  is 86% and  $\text{Ti}^{3+}$  is 14% for iron-doped  $\text{TiO}_2$ . However, mixed  $\text{TiO}_2$  consisted of 93% of  $\text{Ti}^{4+}$  and 7% of  $\text{Ti}^{3+}$ . Similarly in the O1s spectra,  $\text{TiO}_2$  revealed 90% of lattice oxygen and 10% of  $\text{Ti}_2\text{O}_3$ . At the same time iron-doped  $\text{TiO}_2$  consisted 91% of lattice oxygen and 9% of  $\text{Ti}_2\text{O}_3$ . However, mixed  $\text{TiO}_2$  have 97% of lattice oxygen and 3% of  $\text{Ti}_2\text{O}_3$ . The results reveal that both iron-doped and rutile  $\text{TiO}_2$  have comparable oxygen vacancies or defects and mixed  $\text{TiO}_2$  has the least. Besides, iron dopant on the surface of  $\text{TiO}_2$  plays a role as electron carrier which will help create more  $\text{Ti}^{3+}$  and meanwhile help nitrogen adsorbed on the

surface of TiO<sub>2</sub> easier<sup>29,30</sup>. Therefore the results are consistent to the order of photocatalytic testing with iron-doped TiO<sub>2</sub> > rutile TiO<sub>2</sub> > mixed TiO<sub>2</sub>.

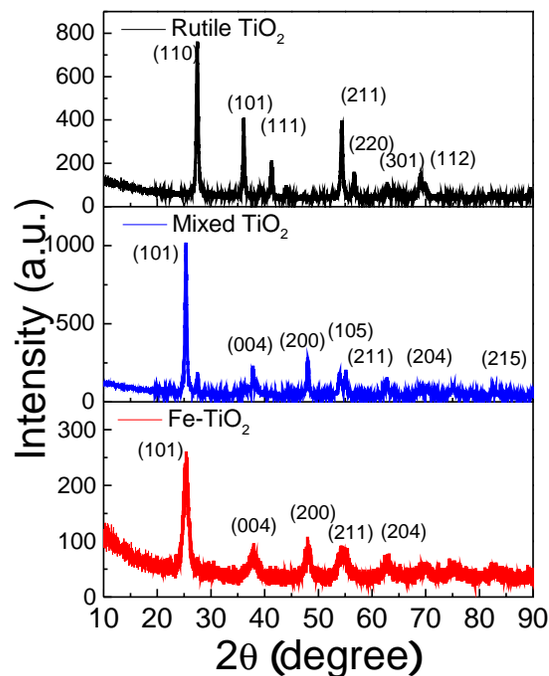


**Fig. 4** XPS of rutile, mixed phase and iron-doped titanium dioxide (a) Ti2p and (b) O1s regions.

In addition to the surface properties, the XRD pattern was recorded to identify the structure of three catalysts. All diffraction lines are relatively strong indicating a high crystallinity for all the samples. Furthermore, the peak positions and relative intensities of the diffraction lines match well with the standard diffraction data for different titanium dioxide phases (i.e. rutile, anatase). For rutile titanium oxide, the major peaks observed at  $2\theta$  values of 27.4°, 36.1°, 41.3°, 54.4°, 56.3°, 69.3° and 70.1° could be indexed to (110), (101), (111), (211), (220), (301), (112) (Figure 5). The pattern is consistent to the standard XRD data (JCPDS, no. 21-1276). The pattern of mixed titanium dioxide is consistent to the XRD standard data as (JCPDS, no. 21-1272 which showed strong anatase crystalline.<sup>31,32</sup> Besides, the XRD spectra of synthesized iron-doped titanium dioxide revealed to be more likely to anatase rather than rutile.

## 4 Conclusions

Photocatalytic nitrogen fixation on titanium dioxide has been heavily investigated for nearly eighty years due to the potential relevance toward achieving the long standing goal of clean ammonia synthesis. Furthermore, nitrogen photofixation is also an intriguing entry point to the abiotic nitrogen cycle, yet has not been thoroughly investigated due to a lack of understanding regarding how the catalytic process occurs on this wide-band gap material. Here we aimed to test an emerging hypothesis which



**Fig. 5** XRD analysis of the rutile, mixed phase and Fe-doped titanium dioxide based photocatalyst

emphasizes that carbon species may play an integral role in catalyzing the dinitrogen adsorption and ultimately reduction to ammonia. Through a controlled experimental procedure whereby methanol is oxidized to produce adsorbed carbon radical based species, we show that the rate of ammonia produced can be increased by as much as two times that observed through testing without adsorbed carbonaceous species. This work extends previous work conducted in the gas phase, which also highlighted the potential role between adventitious carbon and gas phase dinitrogen. Through this work, we also investigate a method to detect ammonia at low levels using a rotating ring disk electrode experimental set up. The results show that a noticeable ammonia oxidation peak is observed on the rutile and iron-doped titanium dioxide samples, yet is not detected with the mixed phase titanium. With many questions regarding the robust nature of low level ammonia measurements, the potential to use current rather than optical or chromatography based methods could improve the resolution of micromolar measurements which is important for both photo and electrochemical nitrogen fixation based studies.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The researchers thank Georgia Tech start up funding for supporting this work, as well as the Georgia Tech TI:GER program.

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