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Photocatalytic oxidation of NOx under visible light on asphalt pavement surface --Manuscript Draft--

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Abstract:	This examines the potential of heterogeneous photocatalysis, as an innovative oxidation technology. The aim is to demonstrate that this technology, is able to reduce the damaging effects of vehicle emissions by using nitrogen doped (N-doped) TiO2 as a photocatalyst immobilized above the asphalt road surface. In the study, the photocatalytic effectiveness and durability of N-doped TiO2 photocatalytic asphalt road material were assessed in both the laboratory and the field by direct and indirect measurement. The experimental results show that N-doped TiO2 asphalt road material has a higher activity compared with the pure TiO2 asphalt road material under visible light irradiation, with decontamination rate for NOx about 27.6%, 24.6%, 16.3% and 13.8% under the irradiation of light wavelength 330-420 nm, 430-530 nm, 470-570 nm and 590-680 nm, respectively. Results of the field test and predicted models suggest that the service life of N-doped TiO2 asphalt road material is approximately 13 months.					
Corresponding Author:	Xiurong Guo Northeast Forestry University haerbin, CHINA					
Corresponding Author E-Mail:	785199452@qq.com					
Order of Authors:	meng chen					
	David Baglee					
	Jiang Wei Chu					
	Danfeng Du					
	Xiurong Guo					
Suggested Reviewers:	yanhua liu liuyanhua815@126.com she is an specialist on the area.					
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Papers published in ASCE Journals must make a contribution to the core body of knowledge and to the advancement of the field. Authors must consider how their new knowledge and/or innovations add value to the state of the art and/or state of the practice. Please outline the specific contributions of this research in the comments box.	 The manuscript utilized asphalt road material and N-doped TiO2 to make environmental protection materials that has higher deNOx ability under the irradiation of visible light. The manuscript has assessed the effectiveness of nitrogen oxides (NOx) degradation by the environmental protection materials under different light wavelength and analysed purification mechanism. Field test and predicted models suggest that the service life of environmental protection materials is for a max period approximately 13 months.
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- Photocatalytic oxidation of NO_x under visible light on asphalt pavement surface 1
- 2 Meng Chen¹, David.Baglee², Jiang Wei Chu³, Dan feng Du⁴, Xiu rong Guo⁵
- 3 ¹College of Traffic, Northeast Forestry University, 26 Hexing Road, Xiangfang District, Harbin,
- 4 150040, P.R. China, E-mail: chenmeng623@126.com
- ² Faculty of Applied Sciences, University of Sunderland, ST PETERS WAY, Sunderland, SR6 5
- 6 0DD, United Kingdom, E-mail: David.Baglee@sunderland.ac.uk
- 7 ³College of Traffic, Northeast Forestry University, 26 Hexing Road, Xiangfang District, Harbin,
- 8 150040, P.R. China, E-mail: wildlife_deer@126.com
- 9 ⁴ College of Traffic, Northeast Forestry University, 26 Hexing Road, Xiangfang District, Harbin,
- 10 150040, P.R. China, E-mail: chenyuqi0831@163.com
- ⁵ College of Mechanical and Electrical engineering, Northeast Forestry University, 26 Hexing 11
- 12 Road, Xiangfang District, Harbin, 150040, P.R. China, E-mail: 785199452@qq.com
- 13 Corresponding author: Xiu rong Guo

15 Abstract:

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16 This work examines the potential use of heterogeneous photocatalysis as an innovative oxidation

technology. The aim is to demonstrate that this technology cand reduce the damaging effects of

vehicle emissions by using nitrogen doped (N-doped) TiO₂ as a photocatalyst immobilized above

the asphalt road surface. In the study, the photocatalytic effectiveness and durability of N-doped

TiO₂ photocatalytic asphalt road material were assessed in both the laboratory and the field using

direct and indirect measurement. The experimental results show that under visible light irradiation,

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23 material. The decontamination rate for NO_x is about 27.6%, 24.6%, 16.3%, and 13.8% under irradiation at light wavelengths of 330-420 nm, 430-530 nm, 470-570 nm and 590-680 nm,

respectively. Results of the field test and prediction models suggest that the service life of

N-doped TiO₂ asphalt road material is approximately 13 months.

Key word: Photocatalyst; N-doping; visible light; Photocatalytic asphalt pavement; Nitrogen

29 oxides

Introduction

Urban areas experience high levels of traffic exhaust, which contribute to air pollution, which is a major global concern (Ishihara et al. 2010). Many governments have introduced emission reduction systems in order to decrease emissions. In spite of these efforts, traffic has continued to increase, adding to concerns about the influence of traffic emissions on health and the environment. Researchers have found that asthma is associated with nitrogen oxide (NO_x), which pedestrians are exposed to due in part to the proximity of roads to walkways. Motor vehicle emissions must be decreased in order to reduce nitrogen oxide emissions, and thus reduce asthma rates (Tarek Mohamed et al. 2009). There are a number of methods that can be used to eliminate NO_x and other pollutants.

Recently, heterogeneous photocatalysis has emerged as an ecological technique for controlling air pollutants. In this technique, TiO₂ and pavement materials (such as cement and asphalt) are used together as a photocatalyst; this method has been found to be a promising and valid technology for NO_x control. The de-polluting properties of TiO₂ photocatalytic materials have been assessed by applying this method both in real-world use and in laboratory simulations

performed under different experimental conditions. Pone and Cheung (2006) evaluated the NO removal paving blocks produced by waste materials and TiO₂. Their study included an optimum mix design incorporating recycled sand, glass, 10% TiO₂, and cement achieved 4.01 mgh⁻¹m⁻² NO removal. Husken et al., (2007) performed a comparative analysis of different photocatalytic cementitious products in an optimum laboratory conditions. They found that the efficiency of NO_x degradation varied significantly, with some products achieving 40% degradation while others had no influence. In Bergamo, Italy in 2006 (Guerrini and Peccati. 2007) a 12,000 m² area, was developed on the sidewalk and the road using active paving blocks. Environmental monitoring showed an average NO_x abatement of 45% during the daytime (09:00 -- 17:00).

In asphalt roads, based on the porous characteristics of asphalt road materials, Meng Chen et al. (2010) used permeability technology to apply asphalt nano-TiO₂ as an environmental protection material. This test showed that this type of photochemical catalysis environmental protection material had a purification function and the ability to protect the environment. Marwa Hassan et al. (2013) used the pray method to make photocatalytic asphalt pavements. Laboratory evaluation showed that the maximum NO_x removal efficiency was reached at an application rate of 0.05 L/m². A research team in Italy used a mixed method approach to develop environmental protection materials. TiO₂ was added into asphalt pavements as an apparent layer that is sprayed onto existing pavements. The decrease in efficiency was dependent on the type of TiO₂ nanoparticles used, and the NO_x decrease efficiency ranged from 20–57% (Venturini et al. 2009).

Photocatalytic asphalt pavements mainly use anatase phase TiO₂. The TiO₂ band gap is 3.2eV, which corresponds to wavelengths less than 388 nm. This limits the photocatalytic practice in the UV light region, which amounts to 4% of the solar spectrum, while the key part (45%) falls under

the visible light region (Chun-Hung et al. 2010). Few studies have attempted to use photocatalytic asphalt pavements under visible light irradiation. Therefore, this study focuses on making a photocatalytic asphalt pavement material. As an innovative oxidation technology, it is able to reduce the damaging effects of vehicle emissions by using N-doped TiO₂ as a photocatalyst that is immobilized above the asphalt road's surface under visible light irradiation. The decontamination effect and application durability of the photocatalytic asphalt pavement material are also analyzed in this study.

Experimental

Photocatalysts preparation

A non-metal doping approach (N-doped TiO_2) was used to extend the utilization of the visible region in the solar spectrum. Titanium powder was prepared using a sol-gel approach that used tetrabutyl titanate (TNBT) and distilled water as the titanium precursor and hydrolyzing agent. First, TNBT was mixed with ethanol and distilled water. Then, the mixed solution was stirred using a machine under an 85° C temperature bath for 6 hours. The slurry was dried at 80° C and calcined at 400° C for one hour. Finally, a white powder was obtained, which was pure TiO_2 (see Figure 1).

The pure TiO_2 powder was mixed with urea at molar ratios of 2:1, and then ground in an agate mortar for homogeneity. The mixed powders were heated in a muffle furnace at 500 °C for 3 h, resulting in N-doped TiO_2 (Figure 1).

Figure 1 shows that the N-doped sample appears yellow in color, compared to the white color of the pure TiO₂. Giacomo Barolo (2012) and Shinri Sato (1986) also observed these differences

in color and found that the yellow-colored N-TiO₂ had greater photocatalytic activity.

Preparation of asphalt road material through the addition of N-doped TiO₂ nanoparticles

We performed a series of penetrating load tests in order to attain a photocatalytic asphalt road material sample. First, based on our preliminary research (Meng and Yan hua. 2010), the penetrant and modified N-doped TiO₂ with silane coupling reagent were prepared. Then, the solution of mixed penetrant and modified N-doped TiO₂ was sprayed onto the bituminous sample's surface. In the load process, every sample was sprayed three times by an atmospheric air compressor. The caliber of the air compressor was 1.5mm and the air pressure was 3.0-3.5MPa. The samples were sprayed at a distance of approximately 20 cm and the speed of the aqueous solution jet was set at 8-10g/s.

Figure 2 and 3 display the schemes of the asphalt road material sample before and after spraying.

Analytical methods

Material property analytical methods

A scanning electron microscopy (SEM), and transmission electron microscopy (TEM) were used in this research in order to analyze the material properties of the N-doped TiO₂ and the photocatalytic asphalt road material. The SEM was a commercial Hitachi S2300 instrument with a tungsten hairpin filament. An accelerating voltage of 25 keV was used on gold samples to eliminate charging. Transmission electron microscopy analyses were performed using a JEM-2010 (JEOL) operating at 160 kV. T.

Photocatalytic degradation analytical methods

Direct and indirect measurement strategies were used in this research to evaluate the pollutant removal efficiency. Direct measurements were used in laboratory tests. In the laboratory test, 500 ppb of NO_x was put into the measurement system. After flowing through the photocatalytic asphalt road material, it was exported via the purification examination photoreactor of the system. Measurement results of the inlet and outlet of photoreactor show that the total NO_x concentration, the NO conversion, NO₂ conversion, and NO_x conversion were:

$$NO_{x} = NO + NO_{2}$$
 (1)

$$NO_{conversion} = \left(\frac{C_{NOin} - C_{NOlight}}{C_{NOin}} - \frac{C_{NObin} - C_{NOsb}}{C_{NObin}}\right) \times 100\%$$
 (2)

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$$NO_{2conversion} = \left(\frac{C_{NO_{2}in} - C_{NO_{2}light}}{C_{NO_{2}in}} - \frac{C_{NO_{2}bin} - C_{NO_{2}sb}}{C_{NO_{2}bin}}\right) \times 100\%$$
 (3)

$$120 \qquad NO_{\text{Xconversion}} = \left(\frac{C_{\text{NO}_{\text{X}}\text{in}} - C_{\text{NO}_{\text{X}}\text{light}}}{C_{\text{NO}_{\text{Y}}\text{in}}} - \frac{C_{\text{NO}_{\text{X}}\text{bin}} - C_{\text{NO}_{\text{X}}\text{sb}}}{C_{\text{NO}_{\text{Y}}\text{bin}}}\right) \times 100\% \tag{4}$$

122 The parameter meaning of equations 1,2, 3, and 4 are described in table 1.

The concentration of nitrates serves as evidence of a photocatalytic decrease of NO_x . Nitrates that accumulate on the pavement surface were measured by dissolving them in deionized water. Using the Japanese industrial standard (JIS TR Z 0018 "Photocatalytic materials— air purification test procedure"), the nitrogen compound eluted from the test piece was calculated using the following formula (David et al. 2014):

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$$Q_{w} = Q_{w1} + Q_{w2} = V_{w1} [(NO_{3}^{-})_{w1} / 62] + [(NO_{3}^{-})_{w2} / 62]$$
 (5)

where Q_w = nitrogen compound eluted from the test piece (µmol); V_w = volume of collected washing (mL); NO_3^- = nitrate ion centration eluent from the test piece (mg/L); and W_1 and W_2 = the first and second DI washes, respectively.

Experimental testing: testing system and set-up

Experimental testing system

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Photocatalytic degradation of NO_x was carried out using a continuous flow system that included a gas supply subsystem, photoreactor, and the analytic subsystem. NOx, N2, and humidified air were supplied in the gas supply subsystem. Humidified air was prepared by bubbling air through a gas wash bottle containing water. The desired water vapor level can be obtained by varying the flow rate of the humidified air stream. To obtain a stable NO_x concentration, NO_x is mixed with humidified air in a gas mixer. During the experiments, gas flow rates are controlled by calibrated flow meters. The photoreactor in the flow system is made of quartz glass in accordance with the American Material Test Association standard (ASTMD 5116-1990) and the Japanese industrial standard (photochemical catalysis material-air purification performance test method (Marwa et al. 2010)). To simulate on-road automobile exhaust, the photoreactor is divided into three parts (1) an inlet, (2) purification region and (3) outlet. The inlet section is nearly a cylinder; a fan controls the stream gas velocity. The purification section is rectangular, which simulates the road surface. Humidity and temperature sensors and a heater were installed in the photoreactor, and the photoreactor was irradiated by a simulation light source. Figure 4 shows the entire photoreactor structure. In the analytic subsystem, NO, NO₂, and NO_x concentration is measured by the analyzer.

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Testing set-up

Photocatalytic activity rests: Previous studies on photocatalytic asphalt roads have shown that illumination by a light source is an essential condition for photodegradation of NO_x. Light source

illumination can cause the formation of electron–hole pairs on the photocatalytic asphalt road material and the electron–hole pair purified NO_x (Meng and Yan hua. 2010). In this study, we attempt to increase insight into the reduction of NO_x under photocatalytic asphalt road material by looking at the influence of the light source on the process.

During the serial experiment, the photocatalytic activity of N-doped TiO_2 photocatalytic asphalt road material was investigated by assessing the NO_x decomposition in the photoreactor. The experiment consisted of three steps.

Step 1: the sample was kept in the dark for 40 min, in order to achieve adsorption and desorption equilibrium of NO_x gas in the photoreactor.

Step 2: the light source was switched on. The range of the light wavelength is 330–420 nm, 430–530 nm (blue LED), 470–570 nm (green LED), and 590–680 nm (red LED).

Step 3: Photocatalytic characterizations of samples were shown using different NO_x concentration or decontamination rates. Direct measurement was used throughout the experiment.

Tests of application durability: In order to measure long term effects and examine decontamination durability, it is necessary to apply N-doped TiO₂ photocatalytic to asphalt road material to decontaminate exhaust from traffic in an outdoor environment. Vehicular activity and rainwash are major influencing factors in actual outdoor traffic environments.

A field and simulation test were adopted in order to predict the durability of the N-doped TiO_2 asphalt road material. In the field test, the test roads were washed once per week. At the same time, traffic volume, temperature, wind direction, wind speed, and humidity were recorded.

During the test, the UV–vis spectroscopy (TU-1901) was used to determine absorbency of the gathered sample by N-(1-naphtyl)-cthylcncdiaminc dihydrochloride colori metric (Meng et al.

2014). The entire test was executed under nature flow conditions in the daytime and separate specimen exams lasted one hour. The field test location is shown in Figure 5.

Two separate simulation tests were conducted. First, in order to study the effect of vehicular activity, the Loaded-Wheel Tester was used to simulate loading and wear on the N-doped TiO_2 photocatalytic asphalt road surface (see Figure 6). The photocatalytic activities of samples that were subject to wear were investigated by evaluating the decomposition of NO_x in the photoreactor. The second test studied the rain wash effect. The samples were placed in to **a** traffic environment from June to August in Harbin, China. After this period a wash test was conducted. The samples were washed once a week and direct and indirect measurements were used to evaluate decontamination ability.

Results and discussion

Physical properties

TEM was executed to investigate the micrographs and dispersion of N-doped TiO₂ in the penetrant. Fig.6 shows that the micrographs of the N-doped TiO₂ are spherical and there is no significant shape change. The findings show that urea can help to hold or diminish the mean size and increase the homogeneity of the size distribution via the synthesis method, which has been described by other researchers (Hao-Hong et al. 2013). N-doped TiO₂ was also well dispersed in the penetrant as seen in Figure 8. An integrated method was used to ensure the dispersion effect of N-doped TiO₂ in the penetrant. It included three techniques. One technique used a silane coupling agent as a modifier, which should help N-doped TiO₂ to have good compatibility in the penetration process.

N-doped TiO₂. In the third tenchnique, an ultrasonoscope was used to disperse the mixing solution in order to reduce spontaneous reunion and flocculation.

The comprehensive dispersion method achieved a good dispersion effect for the following reasons: (1) The affinity between the nanoparticles and the solvent increased; meanwhile, it was easier to open the nano-granular equipment, reducing the dispersion time and energy consumption.

(2) The strong turbulent motion of the liquid flow caused by the magnetic stirring broke and suspended the nano-particle and the ultrasonic agitation continued to disperse the nanoparticles into small particles, which led to a wide range of the solution region. (3) In the modifier component, one end of the active groups can be adsorbed on the surface of the crashed nanoparticles and the other end of the solvent formed adsorption layer, which produced electric charge repulsion, guaranteeing long dispersed suspension of the nanoparticles in the solvent, avoiding flocculation and ensuring the stability of the system.

Photocatalytic activity and purification mechanism

Figure. 9 shows the decontamination ability of the pure TiO₂ asphalt road material and N-doped TiO₂ asphalt road material, where the different wavelength lights (330–420 nm, 430–530 nm, 470–570 nm, and 590–680 nm) were irradiated to the samples. As seen in Figure 8, there was no loss of photoactivity in 330–420 nm wavelength lights (including in the UV section) on the pure TiO₂ asphalt road material and N-doped TiO₂ asphalt road material. Meanwhile, the N-doped TiO₂ asphalt road material had NO_x decontamination abilities of about 27.6%, 24.6%, 16.3%, and 13.8% under the irradiation of light wavelengths 330–420 nm, 430–530 nm, 470–570 nm, and 590–680 nm respectively. Further, on the N-doped TiO₂ asphalt road material under light irradiation, the NO decontamination ability was better than the NO₂ decontamination ability.

220 A number of results can be extracted from Figure 9.

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(1) N-doped TiO₂ asphalt road materials have better photocatalytic activity under visible light 222 There are two potential reasons for this. First, it is well known that photocatalysts need light 223 illumination to trigger photocatalytic oxidation. Figure 8 shows that the N-doped TiO2 asphalt road material had better NO_x decontamination ability under a wavelength range of visible light. 225 The N-doped TiO₂ asphalt road material absorbed visible light to trigger photocatalytic oxidation. 226 Second, the behavior of the N-doped TiO₂ asphalt road material in the NO_x oxidation was also associated with the active species (•OH and O2 •-) formed on the photocatalyst's surface 228 (Todorova et al. 2013). Excite electrons were required from the valence band to create electron 229 and hole pairs. Equation 6 (Shu et al. 2004) shows that energy decreases from irradiation for 230 exciting surface electrons and holes with increasing wavelength of the visible light. It indicated that the band gap might be significantly narrowed by mechanical chemistry doping of nitrogen in 232 titanium, for instance 3.08 eV (Gagoa et al. 2012).

$$E_{g} = 1239.8 / \lambda$$
 (6)

- where E_g is the band gap (eV) of the sample, λ (nm) is the wavelength of the onset of the 234 235 spectrum.
- 236 (2) N-doped TiO₂ asphalt road material had different photocatalytic activity under different 237 visible light wavelengths.
 - Figure 9 shows that the activity for the N-doped TiO₂ asphalt road material was clearly better than the pure TiO2 asphalt road material activity with increasing wavelengths. The findings were attributed to the introduction of N in the TiO2 lattice and the creation of extra electronic states in the TiO₂ band gap (Kumar et al. 2011).

The N-doped TiO₂ samples possessed a two-step adsorption spectra. The first step may have been related to the original titanium band structure and the second adsorption step (in the visible light region) may have been related to the formation of additional electronic states during nitrogen doping (Shu et al. 2008).

Figure 8 shows that decontamination of the NO_x ability of the N-doped TiO₂ asphalt road material decreased with increasing wavelength of the light in. The rate of electron and hole pair recombination was a key factor affecting the decontamination NO_x ability of N-doped TiO₂ asphalt road material (Yao-Hsuan et al.2011). Photoluminescence emission occurs when wavelength photoenergy is absorbed to excite an electron from the valence band and then a longer wavelength luminescence was emitted via recombination of the electron-hole pair (Chun-Hung et al. 2012). The greater the luminescence intensity was, the quicker the electron and hole recombined. Therefore, high intensity of luminescence might result in low photocatalytic activity (Yu et al., 2005). Therefore, the N-doped TiO₂ is capable of lowering the luminescence intensity and guiding the higher photocatalytic activity under areas with shorter wavelengths.

(3) N-doped TiO₂ asphalt road material had a complicated photocatalytic mechanism.

By following the experimental results of the present study, we found that NO₂ decontamination ability was lower on the N-doped TiO₂ asphalt road material under light irradiation. It has been shown that NO₂ was created in the photocatalytic purification process to decrease NO₂decontamination ability. This may be explained by a tentative mechanism that is proposed to explain photocatalytic oxidation of NO_x under visible light on the N-doped TiO₂ asphalt road surface, which is shown in Figure. 10.

It is widely recognized that photocatalytic oxidation can be divided into 3 core stages: first

the transfer of contaminants from bulk to the surface; second the adsorption on the catalyst surface and formation of reactive ions; finally, degradation by the ions formed on the surface (Chuck et al.2 013). Based on Figure 10, a possible photocatalytic NO_x oxidation mechanism was proposed using equations (7)-(18).

N doping can build impurity states below the bottom of the conduction band of TiO₂ asphalt material, causing visible TiO₂ asphalt material to be activated by visible light (Hongqi et al.2011).

N-doped TiO₂ asphalt road material was irradiated by appropriate light and energy, creating a hole and electron pair in the N-doped TiO₂ lattice, shown in equation (7).

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$$N-TiO_2 + visible light \rightarrow h_{VB}^+ + e_{CB}^-$$
 (7)

During adsorption of the reactants onto the N-doped TiO_2 asphalt road material, the photogenerated hole in the conduction band will absorb water, oxygen, nitric oxide, and nitrogen dioxide in the air as described in equations (8) - (11).

$$N-TiO_2 + H_2O \Leftrightarrow N-TiO_2 - H_2O$$
 (8)

$$N-TiO_2 + O_2 \Leftrightarrow N-TiO_2 - O_2$$
 (9)

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$$N-TiO_2 + NO \Leftrightarrow N-TiO_2 - NO$$
 (10)

$$N-TiO_2 + NO_2 \Leftrightarrow N-TiO_2-NO_2 \tag{11}$$

In our previous work (Meng et al.2011), we found that adsorbed H_2O on the N-doped TiO_2 leads to the formation of a highly hydroxylated surface and also gives off hydrogen ions (H^+) and hydroxide ions (OH^-) by its dissociation. Then, H^+ reacts with OH^- to generate • OH^- .

Additionally, the photoinduced electron reacts with O_2 , forming the superoxide anion (O_2^-) and O_2^- forms HO_2 • radicals with traces of water. Lastly, nitrogen species (NO, NO₂) in the air can be easily photocatalytically oxidized leading to HNO_3 by HO_2 • radicals on the N-doped TiO_2

asphalt road surface. The reaction is illustrated in equations (12) to (18).

$$H_2O \rightarrow H^+ + OH^- \tag{12}$$

$$h^{+} + OH^{-} \rightarrow \bullet OH \tag{13}$$

$$e^- + O_2 \rightarrow O_2^- \tag{14}$$

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$$HO_2 \bullet + NO \rightarrow NO_2 + \bullet OH$$
 (15)

$$NO + \bullet OH \rightarrow HNO_{2}$$
 (16)

$$4 \text{HNO}_2 + \bullet \text{OH} \rightarrow \text{NO}_2 + \text{H}_2\text{O}$$
(17)

$$NO_2 + \bullet OH \rightarrow HNO_3 \tag{18}$$

Durability Analysis

Figure. 11 shows the downward trend, over consecutive months, of the NO_x decontamination ability on the N-doped TiO_2 asphalt road. In order to predict the durability of the N-doped TiO_2 asphalt road, we hypothesized that the N-doped TiO_2 asphalt road will actively degrade when the NO_x decontamination rate is less 5%. An exponential regression model was used with the observed data that was obtained in the field test (see Figure 11). The model was $y=32.01e^{-0.145x}$, with 95% confidence. Results from the model showed that active degradation of the asphalt pavement lasted for a period of about 13 months. The photocatalytic efficiency defeat was due to the wear from traffic and the rain wash. Wear and wash simulation tests were completed in order to analyze these reasons.

In the wear simulation test, as seen in table 2, the data showed s decrease in NO_x decontamination ability after wearing. However, the decrease was small under a certain scope, such as in 8000 cycles where the wheel applied a load of 600 N for every cycle. But when the cycle increased (exceeding 8000 cycles), the degree of decrease in the NO_x decontamination

ability increased. This had two possible causes: (1) the samples kept higher NO_x removal efficiencies after wear, and, (2) N-doped TiO₂ was retained on the asphalt road surface and interior after wearing, which may be proven by figure 12. This may be attributed to the research team's use of infiltration liquid with N-doped TiO₂ and the permeability loading method. The infiltration liquid had a special lipophilic penetration ability. When the infiltration liquid is sprayed on the road surface, the nano-material (N-doped TiO₂) can be guaranteed to gradually infiltrate the road surface under the action of the penetrating agent molecular power. Subsequently, with the growth of the concentration of the asphalt pavement surface infiltration liquid and the dual influences of the concentration gradient and porous open graded pavement, N-doped TiO₂ penetrates along the open graded asphalt pavement pore to the core under the action of capillary force and gravity. During the penetration process, when the N-doped TiO₂ molecules are near the asphalt mixture solid surface, N-doped TiO₂ molecules can adsorb on the solid surface in order to achieve the load. This is because of the electric dipole moment and is possible because of the help of the Van Der Waals force. However, when the cycle increases, the loss of weight in the samples also increases. Surface wearing and particle loss may be associated with the loss of N-doped TiO₂ particles, which is shown in figure 13. NO_x decontamination ability decreases as the cycles increase. Table 2 presents the average NO_x decontamination ability for both the original and the washed samples. The table shows that washing the samples results in reduction in the NO_x removal efficiency and this decline was aggravated with time. This indicates time dependency, which will result in a decrease of the NO_x removal efficiency. There are many reasons for this. First, the decrease can be clarified by referencing the NO_x purification mechanisms found in section 3.2 of this paper. HNO₃ should be speedily produced and accumulated on the top of the

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N-doped TiO₂ asphalt road surface, and this could inhibit the photocatalytic reactions by building a physical fence. Next, regeneration of the purification ability has proven that washing is a better way to keep the NO_x decontamination ability for N-doped TiO₂ asphalt road material. During washing, the HNO₃ was easily removed from the catalyst surface (as indirect measurement results); it can be argued this may contribute to catalyst regeneration. Water can also, through rehydration, replenish the consumed hydroxyl radicals; therefore, it is able to maintain the photocatalyst activity (Meng et al.2011). However, the continuing decrease of the NO_x removal efficiency also indicated that washing did not totally recover the active N-doped TiO₂ asphalt road material. That may be due to the adsorption of indissolvable materials, such as lipids, on the photocatalyst receptor sites of N-doped TiO₂ asphalt road material. Adsorption of NO_x occurred on the N-doped TiO₂ asphalt road.

Summary and future work

N-doped TiO₂ powders were successfully prepared by sol-gel methods with urea. The modified procedures did not change the crystalline structure or the morphology of the N-doped TiO₂ as compared with initial sol-gel TiO₂. TEM analysis revealed that N-doped TiO₂ powders could be well dispersed in asphalt penetrants. Based on the penetrant, N-doped TiO₂ asphalt road materials were successfully prepared using spray methods.

Meanwhile, the N-doped TiO_2 asphalt road materials presented higher activity on NO_x removal than pure TiO_2 asphalt road materials under the irradiation of visible light. The outcome was ascribed to the photocatalytic mechanism. As described by this mechanism, the nitrogen species onto The iO_2 interfaces decreased the absorption band gap energy in the visible light and hindered the electron hole recombination, which helped to improve oxidation of nitrogen oxides.

The durability of N-doped TiO_2 asphalt road materials was evaluated through field and simulation testing. Results suggested that the durability of N-doped TiO_2 asphalt road materials spanned a period of approximately 13 months. The results demonstrated that the TiO_2 nitrogen doping approach would provide a worthy channel for photocatalytic asphalt road materials of highly visible light induced photocatalytic activity for the practical decontamination NO_x application in the medium-term.

Because the application of N-doped TiO₂ asphalt road materials in demonstrating vehicle emissions is still a relatively new field of study, more research should be conducted prior to field application. Our research team plans to evaluate the potential pollution of nitrates created by the photocatalytic processing in land through plant experiments. We will also attempt to build an evaluation system for N-doped TiO₂ asphalt road materials application for demonstrating vehicle emissions.

Acknowledgments

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References

Giacomo Barolo, Stefano Livraghi, Mario Chiesa, Maria Cristina Paganini, and Elio Giamello.

(2012). "Mechanism of the Photoactivity under Visible Light of N-Doped Titanium

374	Dioxide. Charge Carriers Migration in Irradiated N-TiO2 Investigated by Electron
375	Paramagnetic Resonance. "The Journal of Physical Chemistry C 116:20887–20894.
376	Hao-Hong Chen, Fang Lei, Jing-Tai Zhao, Ying Shi ,Jian-Jun Xie. (2013). "Preparation and
377	photovoltaic properties of N-doped TiO2 nanocrystals in vacuum." Journal of Materials
378	Research 28:468-47.
379	Meng Chen, Jiang-Wei Chu. (2011). "NO _x photocatalytic degradation on active concrete road
380	surface d from experiment to real-scale application." Journal of Cleaner Production 19:
381	1266-1272.
382	Meng Chen, Li sheng Jin, Yanhua Liu, Xiurong Guo, Jiangwei Chu. (2014). "Decomposition of
383	NO in automobile exhaust by plasma-photocatalysis synergy." Environmental Science and
384	Pollution Research 21:1242–1247.
385	Meng Chen, Yan hua Liu. (2010). "NO _x removal from vehicle emissions by functionality surface
386	of asphalt road." Journal of Hazardous Materials 174:375–379.
387	Poon CS, Cheung E (2006). "NO removal efficiency of photocatalytic paving blocks prepared
388	with recycled materials." Construction and Building Materials 21: 1746-53.
389	Husken G, Hunger M, Brouwers H. (2007). "Comparative study on cementitious products
390	containing titanium dioxide as photo-catalyst. " In: Baglioni P, Cassar L, eds. RILEM Int.
391	Symp. On Photocatalysis. Environment and Construction Materials. Italy 147–54.
392	G.L.Guerrini, E.Peccati. (2007). "Photocatalytic cementitious roads for depollution, in: P.Baglioni,
393	L. Cassar (Eds.), Proceedings international RILEM symposium on photocatalysis."
394	Environment and construction materials-TDP 2007, RILEM Publications, Bagneux
395	187–194.

396 Ishihara, H., Koga, H., Kitaoka, T., Wariishi, H., Tomoda, A., Suzuki, R. (2010). "Paper-structured

397 catalyst for catalytic NO_x removal from combustion exhaust gas. "Chemical Engineering

- 398 Journal 65: 208–213.
- 399 Marwa Hassan, Louay N. Mohammad, Somayeh Asadi. (2013). "Sustainable Photocatalytic
- 400 Asphalt Pavements for Mitigation of Nitrogen Oxide and Sulfur Dioxide Vehicle
- Emissions." Journal of Materials in Civil Engineering 3:365-371.
- 402 Chun-Hung Huanga, I-Kai Wanga, Yu-Ming Linb, Yao-Hsuan Tsengc, Chun-Mei Lud. (2010).
- 403 "Visible light photocatalytic degradation of nitric oxides on PtOx-modified TiO2 via
- 404 sol-gel and impregnation method." Journal of Molecular Catalysis A: Chemical
- 405 316:163–170.
- 406 Chun-Hung Huang, Yu-Ming Lin, I-Kai Wang, and Chun-Mei Lu. (2012). "Photocatalytic Activity
- 407 and Characterization of Carbon-Modified Titania for Visible-Light-Active
- 408 Photodegradation of Nitrogen Oxides." International Journal of Photoenergy 2012: 1-13.
- 409 Venturini, L., and Bacchi, M. (2009). "Research, design, and development of a photocatalytic
- 410 asphalt pavement. "Proc., 2nd Int. Conf. on Environmentally Friendly Roads, Road and
- 411 Bridge Research Institute, Warsaw, Poland 1–16.
- 412 Marwa M.Hassan, Heather Dylla, Louay N.Mohammad, Tyson Rupnow. (2010). "Evaluation of
- 413 the durability of titanium dioxide photocatalyst coating for concrete pavement."
- 414 Construction and Building Materials 24:1456–1461.
- 415 Todorova.N, T.Vaimakis, D. Petrakis, S. Hishita, N. Boukos, T. Giannakopoulou, M. Giannouri, S.
- Antiohos, D. Papageorgiou, E. Chaniotakis, C. Trapalis. (2013). "N and N,S-doped TiO₂
- 417 photocatalysts and their activity in NOx oxidation. "Catalysis Today 209:41–46.

- 418 Tarek Mohamed Naser, Isao Kanda, Toshimasa Ohara, Kazuhiko Sakamoto, Shinji Kobayashi,
- 419 Hiroshi Nitta, Taro Nataami. (2009). "Analysis of traffic-related NO_x and EC
- 420 concentrations at various distances from major roads in Japan." Atmospheric Environment
- 421 43: 2379–2390.
- David Osborn, Marwa Hassan, Somayeh Asadi, John R. White. (2014). "Durability Quantification
- of TiO₂ Surface Coating on Concrete and Asphalt Pavements." Journal of Materials in
- 424 Civil Engineering 26:331-337.
- 425 Gagoa.R, A. Redondo-Cubero b, M. Vinnichenkoc, J. Lehmannc, F. Munnikc, F.J. Palomares.
- 426 (2012). "Spectroscopic evidence of NO_x formation and band-gap narrowing in N-doped
- 427 TiO₂ films grown by pulsed magnetron sputtering." Materials Chemistry and Physics
- 428 136:729-736.
- 429 Shinri Sato. (1986). "Photocatalytic activity of NO_x-doped TiO₂ in the visible light region."
- 430 Chemical Physics Letters 123:126-128.
- 431 Kumar.S.C, L.G. Devi. (2011). "Review on modified TiO₂ photocatalysis under UV/visible light:
- 432 selected results and related mechanisms on interfacial charge carrier transfer dynamics."
- Journal of Physical Chemistry A 115:13211–13241.
- Hongqi Sun, Ruh Ullah, Siewhui Chong, Hua Ming Ang, Moses O. Tadé, Shaobin Wang. (2011).
- 435 "Room-light-induced indoor air purification using an efficient Pt/N-TiO₂ photocatalyst."
- 436 Applied Catalysis B: Environmental 108–109:127–133.
- 437 Yao-Hsuan Tseng, Chien-Hung Kuo. (2011). "Photocatalytic degradation of dye and NO_x using
- visible-light-responsive carbon-containing TiO₂." Catalysis Today 174:114–120.
- Yu.Y, J. C. Yu, J. G. Yu. (2005). "Enhancement of photocatalytic activity of mesoporous TiO₂ by

440	using carbon nanotubes." Applied Catalysis A289: 186–196.
441	Shu Yin, Bin Liu, Peilin Zhang, Takeshi Morikawa, Ken-ichi Yamanaka, and Tsugio Sato. (2008).
442	"Photocatalytic Oxidation of NO _x under Visible LED Light Irradiation over
443	Nitrogen-Doped Titania Particles with Iron or Platinum Loading." Joural of Physical
444	Chemistry. C 112: 12425–12431.
445	Shu Yin, Hiroshi Yamaki, Qiwu Zhang, Masakazu Komatsu, Jinshu Wang, Qing Tang, Fumio Saito
446	Tsugio Sato. (2004). "Mechanochemical synthesis of nitrogen-doped titania and its visible
447	light induced NO _x destruction ability." Solid State Ionics 172:205–209.
448	Chuck W.F. Yu, Jeong Tai Kim. (2013). "Photocatalytic Oxidation for Maintenance of Indoor
449	Environmental Quality." Indoor and Built Environment 22: 139-51.
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Table 1 parameter nomenclature

Parameter	Meaning		
NO _{conversion} (%)	effective purifying rate of NO		
NO _{2conversion} (%)	effective purifying rate of NO2		
NOxconversion (%)	effective purifying rate of NO _x		
C _{NOin} (mg/m ³)	the initial steady-state NO concentration (before turn on		
	the light source)		
C_{NO2in} (mg/m ³)	the initial steady-state NO ₂ concentration (before turn		
	on the light source)		
C_{NOxin} (mg/m ³)	the initial steady-state NO_x concentration (before turn		
	on the light source)		
$C_{NOlight}$ (mg/m ³)	the NO concentration during irradiation phase		
C _{NO2light} (mg/m ³)	the NO ₂ concentration during irradiation phase		
C _{NOxlight} (mg/m ³)	the NO_x concentration during irradiation phase		
C_{NObin} (mg/m ³)	the initial steady state NO concentration		
C_{NO2bin} (mg/m ³)	the initial steady state NO2 concentration		
C_{NOxbin} (mg/m ³)	the initial steady state NO _x concentration		
C_{NOsb} (mg/m ³)	the NO concentration at the end of the blank		
	experiment(without irradiation)		
C_{NO2sb} (mg/m ³)	the NO ₂ concentration at the end of the blank		
	experiment(without irradiation)		
$C_{NOxsb}(mg/m^3)$	the NO _x concentration at the end of the blank		
	experiment(without irradiation)		

Table 2 Degradation effect of wear and wash

D NO 137	Number of wear (cycle)				Number of wash (time)					
DeNO _x ability	0	4000	8000	12000	16000	0	2	4	6	8
Direct										
measurement-purification	36.4	34.8	32.1	28.1	22.3	36.4	34.2	32.8	29.3	24.6
rate (%)										
Indirect measurement										
nitrate concentrations s in	8.3	8.1	7.2	5 5	3.8	8.3	7.9	7.3	6.1	4.1
the sample (mg/L)	0.3	0.1	1.2	5.5	3.8	0.3	1.9	1.3	0.1	4.1



Fig 1. Pure TiO₂ and N-doped TiO₂



Fig 2. Asphalt road material sample before spray



Fig 3. N-doped TiO_2 asphalt road material sample

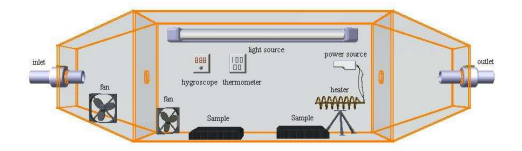


Fig 4. Structure of photoreactor

Fig 5. Spot of field test



Fig 6. Simulation test on vehicular activity effect

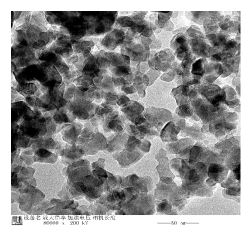


Fig 7. Micrographs of the N-doped TiO₂

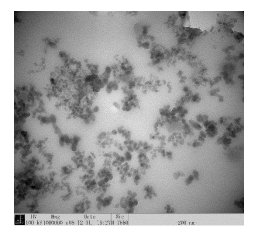
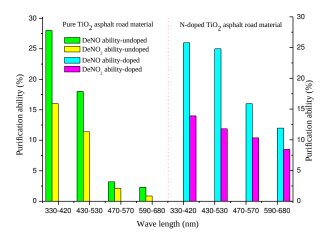


Fig 8. Dispersion effect of N-doped TiO_2 in penetrant



 $Fig \ 9 \ Comparison \ of \ the \ photocatalytic \ activity \ of \ undoped, \ N-doped \ TiO_2 \ asphalt \ road \ material \ under \ UV \ and \\ visible \ light$

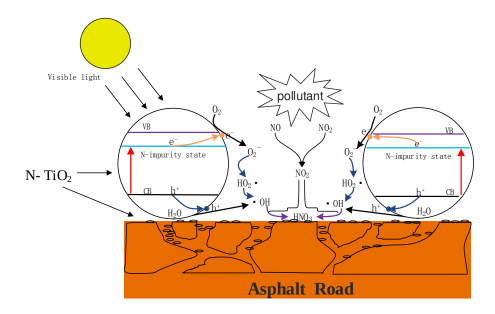


Fig 10. Schematic illustration of photocatalytic process on N-doped TiO2 asphalt road material under visible light

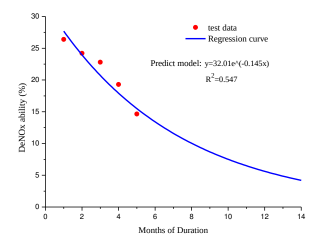


Fig 11 Illustration of durability of N-doped TiO₂ asphalt road

Fig 12 SEM image of road surface after wearing

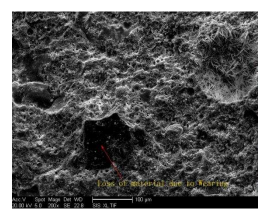


Fig 13 SEM on loss of N-doped TiO_2 due to wearing

Fig	1.	Pure	TiC	\mathbf{e}_2 and	l N-c	loped	TiO ₂

- Fig 2. Asphalt road material sample before spray
- Fig 3. N-doped TiO₂ asphalt road material sample
- Fig 4. Structure of photoreactor
- Fig 5. Spot of field test
- Fig 6. Simulation test on vehicular activity effect
- Fig 7. Micrographs of the N-doped TiO₂
- Fig 8. Dispersion effect of N-doped TiO2 in penetrant
- Fig 9 Comparison of the photocatalytic activity of undoped, N-doped TiO2 asphalt road material under UV and

visible light

- Fig 10. Schematic illustration of photocatalytic process on N-doped TiO₂ asphalt road material under visible light
- Fig 11 Illustration of durability of N-doped TiO₂ asphalt road
- Fig 12 SEM image of road surface after wearing
- Fig 13 SEM on loss of N-doped TiO₂ due to wearing

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Response

Reviewer #2:

Most of my comments have been addressed. However, the paper still needs a revision to improve the language. I would suggest sending to a professional editor before publication.

Answer:

I have found a professional editor to revise the paper.

Response 7: not addressed. The sentence I was referring to is "Indirect measurement based on ..." **Answer:**

The sentence has been deleted.

Response 8: You can make a short statement regarding this issue in the paper.

Answer:

I have made a short statement regarding this issue in the paper. It is added in "Summary and future work". The new expression is "Because the application of N-doped TiO2 asphalt road materials in demonstrating vehicle emissions is still a relatively new field of study, more research should be conducted prior to field application. Our research team plans to evaluate the potential pollution of nitrates created by the photocatalytic processing in land through plant experiments. We will also attempt to build an evaluation system for N-doped TiO2 asphalt road materials application for demonstrating vehicle emissions."