Synthesized and Characterization TiO2-AlLiH4 Nanostructure for Photocatalytic Activity Application for Removal the Residue of Hydrocarbons in Environment

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Abstract

The novel approach includes The sol-gel technique was used to make titanium dioxide nanoparticles and then for photodegradation Benz[a]anthracene a crystalline, aromatic hydrocarbon, as a trial sample of the residue of hydrocarbons in environments close to oil refining facilities and oil fields, methods, the bandgap was modified by solid-state reaction with Lithium aluminium hydride (AlLiH4) reductive compounds. The optical properties were measured using a UV-visible spectrophotometer (Absorbance (A), energy bandgap (Eg), and absorption coefficient (α). TiO2 and TiO2- solid-state reaction showed a clear blue shift of the absorption bandgap which were (2.8ev, 2.7ev, 2.25ev, 2.0ev) to) TiO2, calcination TiO2-AlLiH4 (500C°) then TiO2 -AlLiH4 (750 C°)) respectively. The structure of prepared TiO2 Nanopowders was identified using XRD, Distribution of particle size varied significantly compared to the Sheerer formulation predicted crystallite size (D). which was in a good accordant compared with ASTM results, the particle size and their distribution were characterized using (AFM). To the surface forms and compositions diameters of nanoparticles) SEM) was implemented. The microanalysis of energy scattering X-ray (EDX) was used to examine the chemical makeup of the entire samples. Using UV-Vis spectrophotometry, the photocatalytic response was examined. The photocatalyst impact on the benz[a]anthracene decomposition rate by using catalyst TiO2- and solid-state (TiO2-NaBH4 (550,750)) and (TiO2-AlLiH4 (500,750 C°). A photocatalytic effect is the
catalyst's influence on the catalyst. Using the constant weight of the catalyst and benz[a]anthracene The most effective weight was found (1*10^{-4} M).

**Keywords:** Nanocomposite, Hydrocarbons removal, Photocatalytic, TiO$_2$ - AlLiH$_4$.

1. **Introduction**

Anion doping in broad band gap photo catalysts active under UV irradiation is another way to improve the visible light response [1-3]. Development of visible light photo catalysts from oxide semiconductors by doping with anions as examples (C, N, F, P, and S). The sol-gel technique was used to make AlLiH$_4$-doped TiO$_2$, with different titania precursors and nitrogen sources. Utilizing titanium(IV) tetraisopropoxide with AlLiH$_4$ solution and calcined at 673 K, it was discovered that AlLiH$_4$-doped TiO$_2$ using titanium(IV) tetraisopropoxide with AlLiH$_4$ solution offered the most suitable characteristics for serving as the photo catalyst [5-8]. The TGA, Raman, and XRD findings showed that this AlLiH$_4$-doped TiO$_2$ catalyst had a high crystallinity since the titania precursor had been fully hydrolyzed, with no organic component to obstruct initial phase formation. The results of SEM and TEM showed that the surface shape...
was spherical, similar to fluffy powders [8-11]. Furthermore, the inserted metal or non-metals that’s slowed the anatase-to-rutile phase transition as the calcination temperature rose. The findings of the elemental analysis and UV-Vis/DR showed that ions, metals, and nonmetals may be latent in the TiO2 lattice with strong connections, producing an influence on the band gap structure by adding energy levels near the valence band of TiO2. Under visible light, all of these characteristics improved the photocatalytic activity of co-doped TiO2 [12-15]. In terms of photocatalytic activity, TiO2 - AlLiH4 calcined at 673 K with titanium(IV) tetraisopropoxide and ammonia solution degraded phenanthrene, benz[a]anthracene, and phenol with the best efficiency. When calcined at a higher temperature, however, its photocatalytic activity was substantially reduced. In the instance of photodegradation of phenanthrene, a plausible mechanism was postulated based on two GC/MS identified intermediates: bis(2-ethylhexyl) benzene-1,2-dicarboxylate and dimethyl-4-methyl-1,2- benzene dicarboxylate. [13-22] Incomplete combustion of organic matter produces benz[a]anthracene, a crystalline aromatic hydrocarbon with four fused benzene rings. Benz(a)anthracene can be found in gasoline and diesel exhaust, tobacco and cigarette smoke, coal tar and coal tar pitch, coal combustion emissions, charcoal-broiled foods, amino acids, fatty acids, and carbohydrate pyrolysis products, wood and soot smoke, and creosote, asphalt, and mineral oils, among other things. Only for research purposes is this chemical utilized. It's safe to assume that benz(a)anthracene is a human carcinogen. [23-27] (NCI05)

1.2 NCI Thesaurus (NCIth)

Colorless leaflets or plates, as well as coarse gold powder with a greenish-yellow fluorescence, are found in benz[a]anthracene. It's possible that this substance is carcinogenic. [15,28]

1.3 CAMEO Chemicals

Tetraphene is a four-fused benzene ring polycyclic arene with an angular ortho-fused structure. It is a tetraphenes member and an ortho-fused polycyclic arene. First and foremost, phenanthrene is a PAH made up of three fused benzene rings (Table 1). It is a recognized irritant that photosensitizes skin to light and is present in cigarette smoke.
Phenanthrene is a white powder that is soluble in most organic solvents but insoluble in water. Second, benz[a]anthracene is a PAH [28-30], that has a four-member ring structure, as illustrated in Table 1. It is a natural substance that is created when organic material is incompletely burned. The toxicity of benz[a]anthracene and other PAHs is largely focused at tissues with growing cells [31-33]. Some TiO$_2$ - AlLiH$_4$ will be used in the photocatalytic degradation of benz[a]anthracene. Also investigated were the photocatalytic activity and rate constants of each TiO$_2$ - AlLiH$_4$ and P25 TiO$_2$.

Table 1 The structures and general properties of phenanthrene and benz[a]anthracene.

| Structure | Properties |
|-----------|------------|
|           | Molecular formula | Molecular weight (g/mol) | Melting point (K) | Boiling point (K) |
| ![Structure](image) | C$_{18}$H$_{12}$ | 228.29 | 158 | 438 |

Source: Mackie et al., (2016).

Huang et al. (2008) prepare and calcined TiO$_2$ - AlLiH$_4$ at 593 K using a modified sol-gel hydrothermal process using tetrabutyl titanate as the precursor [34]. XRD research provided strong evidence for TiO$_2$'s heterogeneous crystal lattice structure. The particle size distribution was narrow, with an average particle size of 13.5 nm. The high visible-light absorption of AlLiH$_4$-doped TiO$_2$ was apparent in UV-Vis spectra, which extended beyond 550 nm. The as-prepared sample included Ti-O, N-H, TiO-H, H-O-H, and Ti-NOx groups, and the formation of –Ti-NOx (-Ti-O-N-Ti-) might suggest that N atoms were integrated into the TiO2 crystal lattice, which was compatible with XPS data. The thermal study showed that AlLiH4-doped TiO2 may inhibit the phase change of anatase to rutile, because most TiO2 particles were converted into rutile at 973 K in pure TiO2. From all of these characteristics, the AlLiH4-doped TiO2 (A/R =
4:1) mixed crystal sample had strong visible-light response photocatalytic activity, which was attributed to a synergistic impact among its surface acidity, surface chemistry, and surface doping AlLiH4 atoms in mixed crystals has a chemical impact. The adsorption of cationic organic molecules was increased when the acidic surface was reduced. The doping co atoms increased visible-light absorption, and the mixed crystal effect might have a number of positive impacts on photocatalytic activity.

Ksibi et al, (2008) utilized a titanium(IV) (diisopropoxide)bis(2,4-pentadionate) precursor and various nitrogen and sulfur sources to create N- and/or S-doped TiO2 in a sol-gel process. According to the TGA-DTA statistics, the weight reduction was between 40 and 60 percent. The main homogeneous crystalline phase in the N- and/or S-doped TiO2 samples was anatase, as shown by XRD patterns. The process of absorption [35].

The profile of AlLiH4doped TiO2 moved into visible areas (400-500 nm) when compared to P25 TiO2, owing to band gap narrowing of doped TiO2. Under visible light, the effect of N and/or S doping increased the photocatalytic activity of titania powders as measured by Congo red dye discoloration. Because the crystallinity of anatase was enhanced by S-doping, S-doped TiO2 had the greatest activity among the doped TiO2. Furthermore, sulfur atoms inhibited the anatase to rutile phase transition. Palgrave et al. (2008) used high-resolution XPS to investigate the electrical structure of co-doped TiO2, which was made by annealing single crystal rutile (110) substrates in NH3 at high temperatures [38]. The results indicated that ammonia treatment at 873 K resulted in the integration of AlLiH42p states at the valence band's top without a decrease of Ti4+ to Ti3+. The introduction of AlLiH42p states happened at the same location after annealing at 973 K, but it resulted in surface decrease. The band gap in the valence region photoemission spectra narrowed as a result of the doping. Chen et al. (2007) aimed to study the photochemical reaction of benz[a]anthracene on TiO2 particles [36]. The rising in situ DRIFTS peaks at wavenumbers of 2697 and 3762 cm-1, which correspond to surface hydroxyl properties, led them to the conclusion that this reaction took place on the surface of TiO2 particles. Furthermore, a wide band centered at 3608 cm-1 and a band at 1620 cm-1 diminished with time, indicating that adsorbed
H2O dissolved to produce hydroxyl. The major photoproduct was identified as benz[a]anthracene-7,12-dione based on GC/MS findings [37].

2. Methods

Using titanium tetraisopropoxide and a Sol-Gel technique, nano-TiO2 powder was created (TTIP), according to the [15,16-39].

2.1 Solid State Reaction of TiO2

Different weights of powder TiO2 were mixed with equal proportions of NaBH4 and repeated for AlLiH4. Then, calcination process at different temperatures in the furnace to produce in hydrogen gas by dissociation AlLiH4 and NaBH4 at varies temperatures 500 and 550 according to convert TiO2 by reduction process from anatase to rutile at 600 and 800. The calcination continues for 2h, were the powder appear in black and white colors according to calcination temperature. Then, the ratio (1:1) products (TiO2:NaBH4) and (TiO2:AlLiH4) have been characterization by X-ray diffraction, SEM, EDX, AFM and solid state UV-VIS spectrometer.

2.3 Preparation of benz[a]anthracene solution

25.00 ml of 1000 ppm stock benz[a]anthracene solution in pure methanol solvent was prepared by dissolving 0.0250 g standard benz[a]anthracene with a certain volume in a 25 ml volumetric flask. 500.00 ml of 20 ppm benz[a]anthracene solution was prepared by diluting 10.00 ml of stock benz[a]anthracene solution with distilled water and methanol in the volumetric ratio of 1:3. Concurrently, benz[a]anthracene solution was also sonicated, [40-41]

2.4 The calibration curve of benz[a]anthracene solution

Initially, the series of benz[a]anthracene concentrations, 1, 2, 3, 4 and 5 ppm, were prepared by diluting from 20 ppm of benz[a]anthracene solution (as prepared in section 5.2) with the volume of 0.50, 1.00, 1.50, 2.00 and 2.50 ml, respectively. After that, diluted solution was made up the volume to 10 ml by distilled water and, finally, it was ready to measure fluorescence emission using a Perkin Elmer Lambda 35 luminescence spectrophotometer. [39-41]
2.5 Photo degradation of benz[a]anthracene

Five hours consisting of one hour in dark reaction and four hours in photoreaction were set up for benz[a]anthracene degradation. Under the photoreaction, the sample was collected every 30 min until 90 min of degradation. After that, the sample was continually collected every hour. All of samples were taken by passing the general process as mentioned above. Then, only 1 ml of a sample was picked up using an autopipette from the centrifuged sample and made up the volume by distilled water in a 10 ml volumetric flask. Fluorescent measurement at $\lambda_{em}$ of 527 nm was applied to detect the concentration of each sample. All samples were also conducted in three replicates. [19, 33, 35, 40]

2.6 Luminescence Spectrophotometry

The decreasing concentration of reactants namely benz[a]anthracene was followed by measuring the absorption on a Perkin Elmer Lambda 35 spectrophotometer. To get an obvious spectrum, emission slit width or excitation slit width should be also adjusted to find the appropriate condition. Eventually, for benz[a]anthracene, the optimized condition was $\lambda_{ex}$ at 287.0 nm, with a scan speed of 1000 nm/min, the excitation slit width is 10.0 nm and the emission slit width is 2.5 nm [5,13,33].

3. Result and Discussion

3.1 Photo degradation of benz[a]anthracene

This section made only use of AlLiH$_4$-doped TiO$_2$ using titanium(IV) tetraisopropoxide mixed with AlLiH$_4$ and calcined at 673 K as a photocatalyst due to its most appropriate photocatalytic properties such as crystallinity, phase composition, surface morphology as well as AlLiH$_4$ quantity.
The result as shown in Figure (1) suggests that AlLiH₄-doped TiO₂ could catalyze the photo degradation reaction of benz[a]anthracene with 52% conversion. It degraded with the rate constant of 0.074 h⁻¹ as shown in Table (2). In spite of no results about detected intermediate products, it is assured that the reduction of relative concentration of benz[a]anthracene originated from the AlLiH₄-doped TiO₂ effect. Because, in case of no catalyst, there was simply 8% conversion of benz[a]anthracene this results correspondence with [19, 33, 37, 40-43] AlLiH₄-doped TiO₂ prepared with the variation of titania precursors and AlLiH₄ sources was able to be synthesized by the sol-gel method. From all characterization techniques, it can be implied that the calcination temperature had a significant impact on the crystal structure of AlLiH₄-doped TiO₂. With the increasing temperature, AlLiH₄-doped TiO₂ got the larger crystallite size, the higher crystallinity and Transformation of anatase-to-use phase. The lowest crystallite size, the greatest anatasis and the high number of firmly bound Nitrogen are optimal
for the best photocatalytic activity under visible light for AlLiH₄-doped TiO₂.

Titanium (IV) Tetraisopropoxide as a titanium precursor combined to NH₃ as a nitrogen source is calcined to 674 K compared to all AlLiH₄-doped-TiO₂, and AlLiH₄-doped-TiO₂ seemed to be closely within the scope of the ideal photo catalyst. It provided high crystallinity and spherical surface morphology although its crystallite size is 16 nm. Its phase transformation was retarded by the effect of the strongly-bonded nitrogen in TiO₂ lattice. Moreover, the structural determination did also support that the structure of Titanium (IV) tetraisopropoxide was easy to be hydrolyzed by ammonia or water molecules. With regard to photocatalytic activity, AlLiH₄-doped TiO₂ using titanium(IV) tetraisopropoxide mixed with NH₃ and calcined at 673 K provided the highest %conversion among three substrates; 33% of 20 ppm phenanthrene, 52% of 20 ppm benz[a]anthracene and 4% of 20 ppm phenol. It also rendered the fastest rate of reactions, which was able to be observed from the rate constants; 0.058 h⁻¹ for phenanthrene, 0.074 h⁻¹ for benz[a]anthracene, and 0.036 h⁻¹ for phenol. Interestingly, the elevating temperature had an adverse repercussion on photocatalytic activity.

which resulted from thermal decomposition. Meanwhile, P25 TiO₂ has a good ability to degrade benz[a]anthracene up to 22% conversion with the rate constant of 0.058 h⁻¹ because of synergistic effect between anatase and rutile, like photodegradation of phenanthrene.

| Photocatalyst     | Phenanthrene | Benz[a]anthracene | Phenol |
|-------------------|--------------|-------------------|--------|
|                   | % Conversion | Rate constant (h⁻¹) | % Conversion | Rate constant (h⁻¹) | % Conversion | Rate constant (h⁻¹) |
| AlLiH₄-doped      | 33           | 0.9991 0.058      | 52      | 0.9215 0.074      | 4           | 0.9185 0.0036       |
| P25 TiO₂          | 12           | 0.8093 0.052      | 22      | 0.9227 0.058      | 2           | 0.8737 0.0026       |

*% The conversion has been determined from (I_initial – C_final / I_initial) x 100%.
be a procedure to calculate rate constants are also shown in Appendix.

In addition, the probable mechanism of the photodegradation reaction of phenanthrene was proposed based on the two detectable intermediates. A series of dark reactions with absorbance measurement by UV-Vis spectrophotometry, within different periods with the change of pH and temperature. The results indicate that no degradation of the methylene blue dye although all the factors affecting photocatalytic present except UV-light. At the best weight, the influence of many factors on the organic pollutants decomposition rate was investigated, with pH changes varying from acid to non-neutral and alkaline in acid conditions (4.5, 7.1, and 9.4) as the greatest percentage in benz[a]anthracene breakdown. The effects of change in temperature from (20, 30, 40°C) showed that the breakdown rate increased as the temperature for the MB rose. Energy activation (Ea) has equations as follows: where: k: first-order rate constant (min⁻¹) C0: initial concentration Ct: concentration after time . The rate constant for degradation benz[a]anthracene was calculated from a graph Figure (2 a,b) which is a linear plot between (ln(Ct/C0) versus (t). From slope = -k which equals (-k) according to the equation above are listed and Table (2).

The results in Figure (2) show the increase in temperature (293-313) K will increases the discoloration rate of benz[a]anthracene, and also the rate constant increase explained by the fact adsorption is an endothermic process.

been calculated for respective response by expending the equation of Arrhenius equation between (5.73–11.31) kJ mol⁻¹
It was found that the benz[a]anthracene decolorization was obeyed the first-order rate law. Because interactions depend only on the degradation of benz[a]anthracene to it that the highest concentration which is considered basic material interaction, which can be expressed by Kinetic of Degradation of benz[a]anthracene.

Fig. (2): Plot of time versus. ln (C/C0) using a TiO$_2$ -AlLiH$_4$
                   a- TiO$_2$ -AlLiH$_4$ (500) b- TiO$_2$ -AlLiH$_4$ (750)
4. Conclusion

TiO2-AlLiH4 prepared with the variation of titania precursors and AlLiH4 sources were able to be synthesized by the sol-gel method. From all characterization techniques, it can be implied that the calcination temperature had a significant impact on the crystal structure of TiO2-AlLiH4. With the increasing temperature, TiO2-AlLiH4 got the larger crystallite size, the higher crystallinity, and anatase-to-rutile phase transformation. The smallest crystallite size, the highest anatase crystallinity, and a large amount of strongly-bonded AlLiH4 are the ideal properties of AlLiH4-doped TiO2 for the maximum photocatalytic activity under visible light. Compared to all TiO2-AlLiH4, calcined at 750 K seemed to be closed within the scope of the ideal photocatalyst. It provided high crystallinity and spherical surface morphology although its crystallite size is 16 nm. Concerning photocatalytic activity, TiO2-AlLiH4 using calcined at (500,750 °C). Provided the highest rate conversion among three substrates; 52% of 20 ppm benz[a]anthracene and 4% of 20 ppm phenol. It also rendered the fastest rate of reactions, which was able to be observed from the rate constants; 0.058 h-1 for phenanthrene, 0.074 h-1 for benz[a]anthracene, and 0.036 h-1 for phenol. Interestingly, the elevating temperature had an adverse repercussion on photocatalytic activity. In addition, the probable mechanism of the photodegradation reaction of phenanthrene was proposed based on the two detectable intermediates.
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