

THE EFFECT OF CALCINATION TEMPERATURE TO THE COMPOSITE CHARACTERISTICS OF TiO_2SiO_2 NANOPARTICLE

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Received: 2 July 2020

Revised: 14 October 2020

Accepted: 21 October 2020

ABSTRACT

THE EFFECT OF CALCINATION TEMPERATURE TO THE COMPOSITE CHARACTERISTICS OF TiO_2SiO_2 NANOPARTICLE. Self-cleaning material is a material that utilizes the photocatalytic property to degrade organic and inorganic compounds with the help of UV light. One of the materials that have good photocatalytic property is TiO_2 , the photocatalytic property causes TiO_2 to be amphiphilic: becomes hydrophilic when there is light and becomes hydrophobic when there is no light. The photocatalytic property of TiO_2 can be improved with the addition of buffer material such as SiO_2 . TiO_2SiO_2 nanoparticle was synthesized using a sol-gel method with Titanium (IV) Isopropoxide (TTIP) precursors for TiO_2 and Tetraethyl Orthosilicate (TEOS) precursors for SiO_2 and followed by a variation of calcination temperature of 400 °C, 450 °C, 500 °C, and 550 °C for 2 hours. TiO_2SiO_2 composite was synthesized using composition TiO_2 sol 75% and SiO_2 25%. The result of the synthesis TiO_2SiO_2 composite was characterized by Fourier Transform Infra-Red (FT-IR) instrument to determine the functional groups in the composites and X-Ray Diffraction (XRD) instruments to determine the phase, crystallite size and degree of crystallinity in the composite. The purpose of this research is to synthesized TiO_2SiO_2 nanoparticle as a self-cleaning agent with variation of the calcination temperature, to obtain composite characteristics that can support self cleaning. The self-cleaning ability was based on a produced composite characteristic of TiO_2SiO_2 . The result of FTIR characterization showed that at calcination temperature of 400 °C, 450 °C, 500 °C, and 550 °C there was a Ti-O-Si bond at the peak of 948.91 cm^{-1} , 950.77 cm^{-1} , 941.13 cm^{-1} , 942.13 cm^{-1} . The result of XRD characterization showed that at the temperature of 400°C had the best characteristics, the 75.27% anatase phase and brookite phase 24.72%. Calcinations temperature 400°C had best degree of crystallinity of 91.66%.

Keywords: TiO_2SiO_2 nanoparticles, Sol-gel, Calcination temperature, Self cleaning

ABSTRAK

PENGARUH SUHU KALSINASI TERHADAP KARAKTERISTIK KOMPOSIT NANOPARTIKEL TiO_2SiO_2 . Material self cleaning adalah material yang memanfaatkan sifat fotokatalis untuk mendegradasi senyawa organik dan anorganik dengan bantuan sinar UV. Salah satu material yang memiliki sifat fotokatalis baik yaitu TiO_2 , sifat fotokatalis menyebabkan TiO_2 bersifat ampifilik yaitu hidrofilik saat ada cahaya dan hidrofobik saat tidak ada cahaya. Sifat fotokatalis TiO_2 dapat ditingkatkan dengan menambahkan material penyangga seperti SiO_2 . Sintesis nanopartikel TiO_2SiO_2 menggunakan metode sol-gel dengan prekursor Titanium (IV) Isopropoxide (TTIP) untuk TiO_2 dan prekursor Tetraethyl Orthosilicate (TEOS) untuk SiO_2 , diikuti dengan variasi suhu kalsinasi 400 °C, 450 °C, 500 °C, dan 550 °C selama 2 jam. Komposit TiO_2SiO_2 disintesis dengan perbandingan komposisi sol TiO_2 75% dan SiO_2 25%. Komposit TiO_2SiO_2 yang telah disintesis dikarakterisasi dengan Fourier Transform Infra Red (FTIR) untuk mengetahui gugus fungsional

dalam komposit dan X-Ray Diffraction (XRD) untuk menentukan fasa, ukuran kristal dan derajat kristalinitas komposit. Penelitian ini bertujuan untuk melakukan sintesis nanopartikel TiO₂SiO₂ sebagai agen self cleaning dengan memvariasi suhu kalsinasi, agar diperoleh komposit dengan karakteristik yang dapat mendukung sifat self cleaning. Kemampuan self-cleaning didasarkan pada karakteristik komposit TiO₂SiO₂ yang dihasilkan. Hasil karakterisasi FTIR menunjukkan bahwa pada suhu kalsinasi 400 °C, 450 °C, 500 °C, dan 550 °C terdapat ikatan Ti-O-Si pada puncak 94.91 cm⁻¹, 950.77 cm⁻¹, 941.13 cm⁻¹, 942.13 cm⁻¹. Hasil karakterisasi XRD menunjukkan bahwa pada suhu 400 °C memiliki karakteristik terbaik yaitu fasa anatase 75.27% dan fasa brookite 24.72%. Suhu kalsinasi 400 °C memiliki derajat kristalinitas terbaik yaitu sebesar 91.66%.

Kata kunci: Nanopartikel TiO₂SiO₂, Sol-gel, Suhu kalsinasi, Self cleaning

INTRODUCTION

Nanotechnology is currently developing rapidly in all vital areas of science and technology such as electronics, aviation, defense, medicine and health. This relates to modeling, synthesis, characterization, and application of materials and equipment nanometer scale. The physical, chemical and biological properties of nanoscale differ from the properties of atoms and molecules in large materials. Therefore, it can provides an opportunity to develop a new class of advanced materials that have high-tech applications [1].

One of the nanomaterial is TiO₂ and SiO₂. Nanoparticle composite of TiO₂SiO₂ was synthesized by using sol gel method [2]. Sol gel method has advantages of sustainable homogeneity, can be conducted in room temperature, strengthening interaction between main semiconductor and buffer material. The steps of sol gel process are hydrolysis, condensation, aging, drying, and calcinations [3]. The last step is calcination, a heating process which aim to make photocatalyst's surface structure stronger, moreover organic and inorganic polluter in the catalyst can be eliminated during this process [4]. Calcination temperature has an effect on phase composition, crystals size, degree of crystallinity, and the amount of energy of semiconductor material band gap that was produced. TiO₂ would have change of crystal structure if the calcination temperature increased. The crystals size would decrease with the increasing calcination temperature, thus photocatalytic activity going better in degrading a compound [5].

Calcination temperature affects the type of crystals formed. The anatase crystal phase was formed at a temperature of 400 °C and the growth of anatase crystals increased along with the increase in calcination temperature, at a temperature of 400-600 °C rutile nucleation occurred [6]. At a temperature of 600-900 °C, rutile crystal phase growth occurs and grows slowly as the calcination temperature increases [7]. The range of

UV rays absorbed by rutile is greater, but anatase has a greater photocatalytic activity. This is due to the difference in the energy structure between the two types of crystals, where the conduction band of the anatase crystals is closer to valence band so the reduction strength of the anatase crystals is greater than that of rutile [8]. Calcination temperature has affects on crystal size of TiO₂SiO₂ composites. When the calcination temperature higher, the smaller size of the crystals formed, this is because the diffusion of atoms becomes faster, which accelerates the crystallinity and forms a smaller crystal size. Based on other research results, each calcination temperature variation has a different crystal size, at 250 °C has a crystal size of 50.59 nm, 300 °C has a crystal size 23.64 nm, 400 °C has a crystal size 22.85 nm and 500 °C has a crystal size 20.78 nm [9].

Compounding TiO₂ with SiO₂ was done to increase adsorption ability and decrease recombination, so that photocatalytic activity of TiO₂ can increase and decompose organic polluter [1]. Adsorption activity of TiO₂ can be increased by adding buffer material with large surface area and high porosity [3]. SiO₂ is a catalyst material that stable in high temperature and good mechanical resistance so that photocatalytic ability of TiO₂ can be sustained. SiO₂ has large surface area, it has hydroxyl group on its surface which work for adsorption side [10]. Addition of SiO₂ to TiO₂ can reduce the crystal's size, thus can increase the surface area of TiO₂. The increase of surface area caused the increasing amount of active side, which is hole (h⁺) as an acceptor and electron (e⁻) as a donor absorbed and participate in reduction-oxidation reaction, thus its photocatalytic activity was increasing [11]. SiO₂ can increase adsorption surface by water and hydroxyl group that was adsorbed and photocatalytic activity, because phase transformation from anatase to rutile was hampered by thermal stability and can be increased with titania silica oxide. The addition of SiO₂ to TiO₂ can act as a barrier

and delay of C-C bond on polymer matrix [12]. The combination of TiO_2SiO_2 is expected to produce a material that has good crystal size, phase composition and degree of crystallinity.

In this research, the addition of SiO_2 to TiO_2 made TiO_2SiO_2 composite have three unique characteristics physically and chemically, which are photocatalyst with mix of supporting effect and quantum size effect, acid catalyst that associated with new acid site generation [10]. The effect of calcination temperature also supports the properties of the crystal size, degree of crystallinity, and phase composition of the TiO_2SiO_2 composite. According to the explanation above, the researcher has conducted research about the synthesis of TiO_2SiO_2 using sol gel method with variation of calcination temperature at 400 °C, 450 °C, 500 °C, 550 °C. The temperature is used because the TiO_2SiO_2 composite has the best characteristics among the temperature of 400-550 °C. The characterization of nanoparticle of TiO_2SiO_2 was tested using XRD and FTIR instrument.

EXPERIMENTAL METHOD

Materials and Instruments

Nanoparticle composite of TiO_2SiO_2 was synthesized by using sol gel method. Sol-gel method is an alternative method which is good for the synthesis of silica and titania nanoparticles, it can adjust the geometric structure, degree of crystallinity, phase composition and particle size [13]. The steps of sol gel process are hydrolysis, condensation, aging, drying, and calcination. The use of the sol gel method was chosen because it is relatively easy and can produce homogeneous TiO_2SiO_2 nanocomposites using tetraethoxysilane (TEOS) precursors and TiO_2 nanoparticles and can grow on the surface of TiO_2 nanoparticles [14]. The calcinations temperature affects the final composite characteristics. The materials used in this research are titanium (IV) isopropoxide (TTIP, Sigma Aldrich 98%), tetraethyl orthosilicate (TEOS, Sigma Aldrich 98%) were obtained from China, aquades, ethanol (Merck, 99%), hydrochloric acid (HCl, Merck, 37%), glacial acetic acid were obtained from Indonesia.

The tools used in this study are glass, dropper pipette, beaker, measuring cup, hot plate stirrer, magnetic stirrer, spatula, watch glass, mortar and pestle, analytical balance. Furnace, oven (Binder, Germani), ultrasonicator (Ultrasonic Power 540) were done in anorganic laboratory of Unesa, Fourier Transform Infrared Spectrophotometer

(Shimadzu 8021), X-Ray Diffraction (Shimadzu XRD 6000) were done in material and metallurgy laboratory of ITS. TiO_2SiO_2 composite was characterized by Fourier Transform Infra-Red (FT-IR) instrument to determine the functional groups in the composites and X-Ray Diffraction (XRD) instruments to determine the phase, crystallite size and degree of crystallinity in the composite.

Method and Procedure

TiO_2 sol synthesis

The synthesis of TiO_2 was carried out by sol-gel method. Started with 2 mL of glacial acetic acid as a catalyst, mixed with 5 mL of ethanol p.a as a solvent and 3.2 mL of TTIP precursor was added drop by drop at a time while in ultrasonicator for 15 minutes. The result is a clear, light yellow solution which then let it sit for 24 hours until it became gel.

SiO_2 sol synthesis

The synthesis of SiO_2 was carried out by sol-gel method. Started with adding 5 mL of tetraethyl orthosilicate (TEOS) mixed with 5 mL of ethanol p.a. then put it to stirrer for 10 minutes at 300 rpm. Next, 3-4 drops of 0.1 M HCl solution were put into 10 mL of aquades then stirred homogeneously. After that the mix of HCl solution and aquades were poured into TEOS and ethanol p.a solution and put it to stirrer for five minutes at 300 rpm. Next, the solution was heated at 60 °C for approximately 90 minutes and put it to stirrer at 300 rpm. Let it sit for 24 hours until it became gel.

Composite synthesis of TiO_2SiO_2

The composite synthesis of TiO_2SiO_2 was started by mixing TiO_2 and SiO_2 sol with the composition of 75%. Put it in the stirrer until TiO_2SiO_2 sol became homogeneous. Next, let it sit for 24 hours in room temperature for aging process. Then the solution was dried in oven at 110 °C for 4 days until it crystallized. After that, the crystal of TiO_2SiO_2 composite was mashed until it became powder and calcinated the powder in 400 °C, 450 °C, 500 °C and 550 °C.

Characterization of TiO_2SiO_2 composite using XRD

1 mg of each calcination variation of TiO_2SiO_2 composite powder was taken to be tested and analysed using XRD with 2θ between 3° - 100°.

Characterization of TiO₂SiO₂ composite using FTIR

1 mg of each calcination variation of TiO₂SiO₂ composite powder was taken to be tested and analysed using FTIR.

RESULT AND DISCUSSION

TiO₂ synthesis

The addition of TTIP precursor can speed up the hydrolysis process because TTIP has short alkoxy chain. The shorter the alkoxy chain (M-O-R) then the precursor reactivity is bigger and sol-gel hydrolysis process is faster, thus carbon atom chain in the precursor easily reacted with siloxane group (Si-O-Si) forming Si-O-Ti bond. The longer the ligand (-OR) in the precursor, the slower the hydrolysis rate. It is because of steric hindrance due to the longer R chain, therefore the transfer of ligand (-OR) with hydroxyl group (-OH) is slower [15].

In the synthesis process, ethanol solution acted as solvent. The type of solvent affects the hydrolysis process. Solvents that are often used are methanol and ethanol. Methanol has a shorter C chain than ethanol, so methanol has a greater degree of polarity. More polar solvents cause the hydrolysis process to be faster and more stabilize the Ti-OH group, the more stable the nucleus is, the condensation reaction will spread throughout the core and smaller the crystal size is formed [16].

When TiO₂ sol is irradiated with ultrasonic waves, the solution creates a collision between the constituent particles under high pressure. The ultrasonication time affect the particle size, longer the ultrasonication time the particle size to be more homogeneous and smaller, which ultimately leads to a stable nanoparticle size and less agglomeration [17]. The ultrasonic times of 120 minutes and 180 minutes had crystal sizes of 13.64 nm and 11.84 nm. Caused by the shock wave during ultrasonication separate the agglomeration of particles and a perfect disperse occurs with the addition of a factor as a stabilizer [18]. Based on other research [19], the type of solvent and the time of ultrasonication have an effect on the crystal size and characterization results.

Glacial acetic acid acted as catalyst to speed up the hydrolysis process, both aimed to help the formation process of TiO₂. With the condition inside of ultrasonicator, the solution's components would have

increase of activity with the movement of ultrasonic wave under water, so the micro-mixing process in the solvent will be formed [20]. Condensation process occurred which would produce inorganic compound and formed TiO₂.xH₂O. The aim of aging process is to strengthen the result gel with interpolymer bond.

SiO₂ sol synthesis

This synthesis process, ethanol solution acted as solvent. The type of solvent affects the hydrolysis process. Solvents that are often used are methanol and ethanol. Methanol has a shorter C chain than ethanol, so methanol has a greater degree of polarity. More polar solvents cause the hydrolysis process to be faster and more stabilize the Ti-OH group, the more stable the nucleus is, the condensation reaction will spread throughout the core so that the smaller the crystal size is formed [16]. Glacial acetic acid acted as catalyst to speed up hydrolysis process, aquades helped to hydrolyse TEOS in acidic state, all three aimed to help the formation process of SiO₂. To accelerate the hydrolysis process without eliminating the solvent inside. In gel formation, there was SiO₂ tissue surrounded by pores filled with ethanol and H₂O [8].

TiO₂SiO₂ composite synthesis

TTIP act as the hydrolysis agent and group of Si-O-Si in the polymer can be formed, it can form bond of Ti-O-Si. In condensation process, there was a change from sol to gel. The used 110 °C is to dried and evaporate the solvent leftover in the gel and gel that has cross linking addition in unreacted group of -OH and -OR, thus it can form good solids [15]. The anatase crystal phase was formed at a temperature of 400 °C and the growth of anatase crystals increased along with the increase in calcination temperature, at a temperature of 400-600 °C rutile nucleation occurred [6]. At a temperature of 600-900 °C, rutile crystal phase growth occurs and grows slowly as the calcination temperature increases [7]. Based on research from other journals [9], at a calcination temperature of 250-350 °C the crystallite sizes were 50.59 nm, 42.42 nm and 35.94 nm. At temperatures over 550 °C, the crystal size is not optimal.

TiO₂SiO₂ Composite Phase Characterization

The result of X-ray diffractogram of TiO₂SiO₂ was provided in Figure 1.

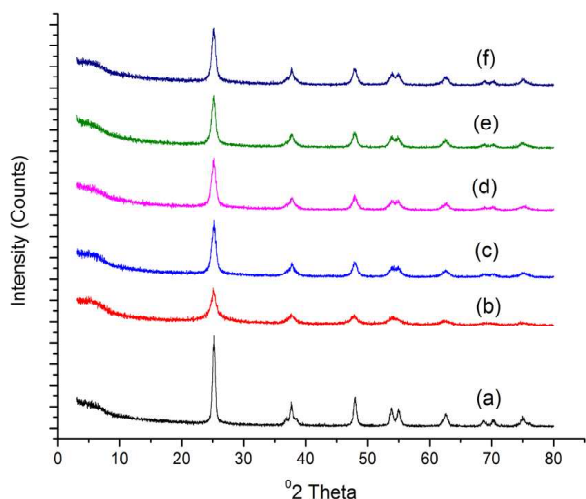


Figure 1. X-ray diffractogram of: (a) standard TiO₂ (b) TiO₂SiO₂ without calcination (c) TiO₂SiO₂ 400 °C (d) TiO₂SiO₂ 450 °C (e) TiO₂SiO₂ 500 °C (f) TiO₂SiO₂ 550 °C

According to XRD diffractogram in Figure 1, it shows that TiO₂SiO₂ composite that was produced has crystalline characteristic, marked with 2θ angle that shows sharp and pointed peaks. Crystalline characteristic is acquired by comparing 2θ value or d(A) in each peak of the XRD diffractogram result using Joint Crystal Powder Diffraction Standard (JCPDS) card. Each calcinations temperature of TiO₂SiO₂ composite shows crystalline characteristic because the ratio of TiO₂SiO₂ is dominated by TiO₂ for 75%.

The addition of SiO₂ into TiO₂ caused SiO₂ enter the gaps of TiO₂ structure and the result is it might break TiO₂ crystal structure. The atoms that formed in the composite during the calcination process was not stable when subjected to heat flow. Those two things made the decrease of intensity and the shift of 2θ to the lower direction [14]. XRD pattern in Figure 1(c) to 1(f) has almost the same peak intensity.

In 400 °C, 450 °C, 500 °C, 500 °C calcination TiO₂SiO₂ composite, the highest peak occurred at the angle of 25.09° followed by peak at 37.80° 53.90° which shows the anatase phase that match with JCPDS number 1272. Other than anatase phase, TiO₂SiO₂ composite with peak at the angle of 47.87° and 68,78° shows brookite phase, it matched with JCPDS number 29-1360. At the angle of 62.74°, it shows the occurrence of rutile phase that match with JCPDS number 21-1276. At the temperature of 400°C, it has anatase phase composition of 75.27%, 54.92% at 450°C, 56.44% at 500 °C, 50.58% at 550 °C. From each of calcination temperature variation shows the more dominant percentage of anatase phase composition than brookite and rutile phase. For more information, composition phase percentage is shown in

Table 1. At 400 °C the optimum crystalline phase of anatase is formed, when the calcination temperature increases 450-550 °C, the anatase phase begins unstable and transforms into rutile. TS has a greater rutile phase composition than standard TiO₂, SiO₂ should inhibit the transformation of TiO₂ from the anatase phase to rutile [21].

According to XRD synthesis result data, it shows that anatase phase of TiO₂SiO₂ composite has bigger composition than standard TiO₂. The composite of TS 400 °C has the biggest anatase phase composition compared to TS, TS 450 °C, TS 500 °C, TS 550 °C. The best formed anatase phase was at the calcination temperature of 400 °C, a higher calcination temperature caused the produced anatase phase to be smaller and starting to be unstable, therefore there is a change of crystal form to rutile and brookite. Rutile phase in TS 550 °C composite was bigger than TS, TS 450 °C, TS 500 °C composite. At a high calcination temperature, SiO₂ moves to gaps of TiO₂ crystal structure thus TiO₂ renovation process was occurred that caused change from anatase phase to rutile phase [9]. Based on mentioned above, TiO₂SiO₂ composite at 400 °C showed the best photocatalytic activity, marked with the biggest anatase phase among every variation of calcination temperature [14].

Characterization of Crystallinity Degree of TiO₂SiO₂ Composite

Result percentage that shows how much crystalline or amorphous content in a material is called degree of crystallinity. According to degree of crystallinity calculation, it was found that 400 °C calcination temperature TiO₂SiO₂ composite has 91.66% degree of crystallinity. At calcination temperature of 450 °C showed degree of crystallinity of 89.74%. At calcination temperature of 500 °C and 550 °C showed degree of crystallinity of 90.90% and 88.70%. For TS and TiO₂ standard sample, they have 83.33% and 90.74% for degree of crystallinity. Those acquired result shows that degree of crystallinity was increasing and decreasing as the calcination temperature got higher. However, degree of crystallinity supposed to be increasing as the calcination temperature got higher [9]. Matching with another research data that state the degree of crystallinity increase as the calcination temperature got higher [22].

Based on the obtained data, sample at calcination temperature of 400°C has the highest degree of crystallinity. It is because of the biggest anatase phase

composition is sample TS 400 °C. Anatase crystal has a good degree of crystallinity because of electron transport in anatase crystal is relatively faster than rutile and brookite crystal [6,7]. A good degree of crystallinity will influence the characteristics of TiO₂SiO₂ composite, the bigger the degree of crystallinity, its photocatalytic ability is also increasing.

Characterization of TiO₂SiO₂ Composite Crystal Size

Based on data of synthesized diffractogram XRD, TiO₂SiO₂ composite crystal's size showed a raise as the increase of calcination temperature. At calcination temperature of 400 °C showed the crystal's size was at 18.30 nm, then decreased to 16.51 nm at 450 °C. Crystal's size at 500 °C and 550 °C was increasing at 19.58 nm and 31.47 nm. The smallest crystal's size was when the composite was at calcination temperature of 450 °C.

Table 1. Data of phase composition and synthesized crystal size

Material	Phase Composition (%)			Crystal Size (nm)
	Anatase	Rutile	Brookite	
TiO ₂ standar	67.89	10.41	21.68	30.86
TS	54.33	13.41	32.25	14.81
TS 400°C	75.27	0	24.72	18.30
TS 450°C	54.92	10.77	34.29	16.51
TS 500°C	56.44	10.93	32.61	19.58
TS 550°C	50.58	18.87	30.54	31.47

According to research result in another journal, higher calcination temperature was supposed to make smaller crystal size, thus the structure of the crystal was more regular and the surface area was getting bigger [8]. At a temperature 550 °C has the largest crystal size. Several things that affect synthesise result are condition influence during synthesise, preparation method, the type of used precursor, and the type of interaction produced between TiO₂ and SiO₂. TTIP is a highly reactive precursor and sustain a rapid hydrolysis process, this cause composite formation with SiO₂ during condensation process was fast [15]. Based on these, it possibly caused agglomeration between neighbouring particles due to fast reaction and increase of particle size distribution that cause the produced crystal's size was getting bigger as the calcination temperature

increase [21]. Mixing process of SiO₂ into TiO₂ was not yet homogeneous, because the produced sol only mixed physically not chemically. That caused the crystal's size getting bigger in high calcination temperature.

Different calcination temperature on TiO₂SiO₂ composite was affecting the characteristics of the produced composite, the higher the calcination temperature the crystal size was getting lower as the decrease of anatase phase. At a temperature 550 °C has the largest crystal size, it possibly caused agglomeration between neighbouring particles due to fast reaction and increase of particle size distribution that cause the produced crystal's size was getting bigger as the calcination temperature increase. The increase of calcination temperature caused the decreasing of composite crystal's size and the surface area is increasing [14].

Characterization of Functional Group of TiO₂SiO₂ Composite Using FTIR

Characterization of functional group of TiO₂SiO₂ composite data is shown in Figure 2.

Based on FTIR result data in Figure 2 (c), (d), (e), (f) shows the peak in wavelength 3217.43 cm⁻¹, 3248.84 cm⁻¹, 3245.92 cm⁻¹, 3254.85 cm⁻¹, 3299.08 cm⁻¹ that shows the occurrence of -OH stretching vibration that caused by hydrogen bond from Si-OH group. Wave number of TS 400 °C composites show absorption area of around 1640-1620 cm⁻¹ that is 1637.08 cm⁻¹. In those absorption areas show the occurrence of -OH bending vibration from hydroxyl because of the absence of hydrogen bond [14]. According to other research result, -OH bending vibration is in absorption area of 1600-1640 cm⁻¹ [23].

Si-O stretching vibration in 400 °C showed the absorption peak in the areas of 1055.97 cm⁻¹. In wave numbers of 422.71 cm⁻¹ showed Ti-O stretching vibration. Those result is almost as the same as research result, which is Si-O bond is in the absorption area of 1050-1090 cm⁻¹ and Ti-O bond is in the absorption area of 420-460 cm⁻¹ [23].

Wave number in TiO₂SiO₂ composite shows absorption area around 950-960 cm⁻¹, based on FTIR result Ti-O-Si bond 400°C formed in absorption areas of 948.91 cm⁻¹. Those areas showed low intensity peaks, it is because of the influence when synthesizing process and its preparation method, thus Ti-O-Si did not perfectly mix chemically (Ti-O-Si bond formation), but perfectly mixed physically (with interaction force not more than Van der Waals weak bond) [6]. As the calcination

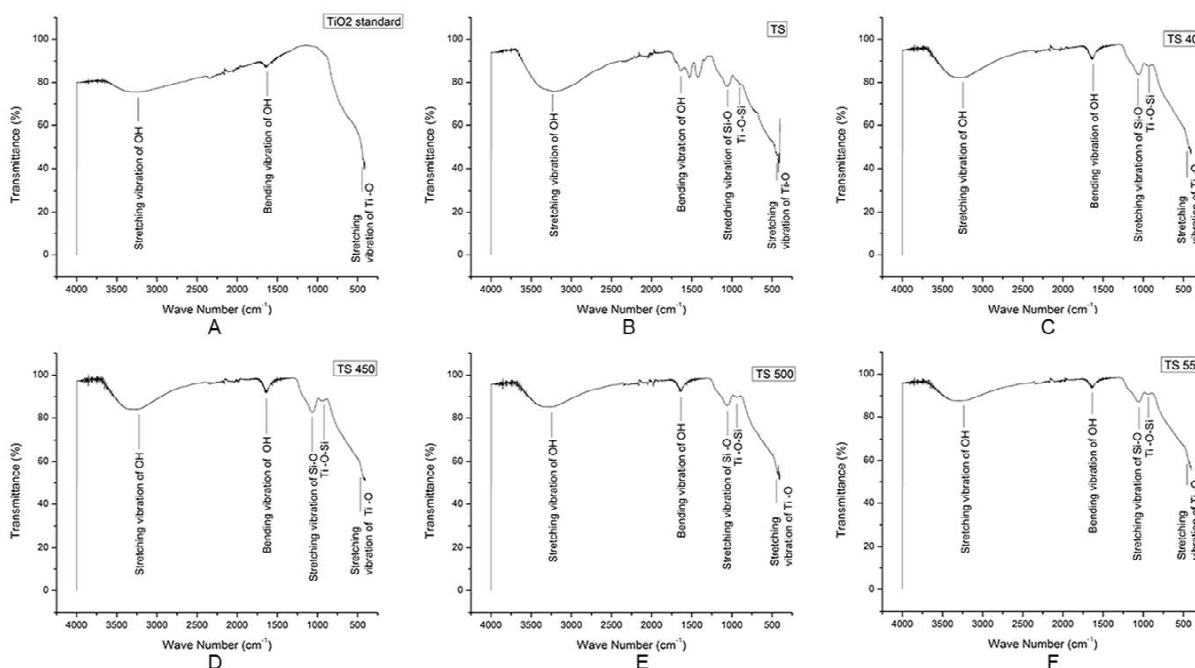


Figure 2. The result of FTIR: (a) TiO₂ standard (b) TiO₂ without calcination (c) TiO₂SiO₂ 400 °C (d) TiO₂SiO₂ 450 °C (e) TiO₂SiO₂ 500 °C (f) TiO₂SiO₂ 550 °C

temperature getting higher, it caused the peaks that consist of bending and stretching vibration of O-H bond getting lower and decreasing its intensity. Table 2 shows absorption areas of wave number that formed form synthesized composite.

Based on Table 2, wave numbers of Ti-O-Si bond at a temperature of 400-550 °C have an insignificant increase and decrease. On higher calcination temperature has lower peak intensity, caused by effect on the ability of TiO₂SiO₂ in self-cleaning activities. Self cleaning activity depends on the peaks of Ti-O-Si bonds are formed. At a temperature of 400 °C, it shows peak intensity reaches the optimum point, because at 450 °C has an increase peak and accompanied by a decrease at temperatures of 500 °C and 550 °C.

According to FTIR result in Figure 2, it shows that synthesized composite showed low peak intensity

of Ti-O-Si. With Si(OC₂H₅)₃(OH) compound being there as a precursor during reaction process can affect crystallinity characteristics of the formed composite, causing SiO₂ to be more amorphous [20]. FTIR result can be strengthen with diffractogram result on 2θ peak that showed that the addition of SiO₂ in the composite can lower crystallinity characteristic of TiO₂SiO₂, thus it seems that TiO₂ peak intensity before the addition of SiO₂ was relatively higher marked with sharper and higher diffraction peak intensity. Based on that, composite with variation difference of calcination temperature does not significantly affect produced functional group.

CONCLUSION

The TiO₂SiO₂ nanocomposites has been synthesized by sol gel method. The characterization of composite showed the effect of calcinations to crystallite

Table 2. Functional group data that formed form synthesized composite

Functional Group	Wave Number (cm ⁻¹)					
	TiO ₂ standard	TiO ₂ -SiO ₂	TS 400°C	TS 450°C	TS 500°C	TS 550°C
Stretching vibration of O-H from Si-OH or Ti-OH	3227.73	3217.43	3248.84	3245.92	3254.85	3299.08
Bending vibration of O-H from Si-OH or Ti-OH	1654.00	1628.50	1637.08	1637.09	1636.98	1637.18
Stretching vibration of Si-O from Si-O-Si	-	1050.48	1055.97	1062.24	1061.87	1056.87
Stretching vibration of Si-O from Si-O-Si	418.83	439.14	422.71	417.30	429.36	424.28
Bonds of Ti-O-Si	-	949.34	948.91	950.77	941.13	942.13

sized, phase composition, degree of crystallinity and wave number. The optimum calcination temperature is 400 °C showed the best characteristic has 75.27% anatase phase composition and 24.72% brookite with crystal's size of 18.30 nm and degree of crystallinity of 91.66%. The presence of TiO₂/SiO₂ composite functional group in wave number of 948.91 cm⁻¹ showed that there is bonds of Ti-O-Si.

ACKNOWLEDGMENT

Thank you for Unesa Instrumental Laboratory, Unesa Inorganic Chemistry Laboratory, ITS Material and Metallurgy Laboratory as a place to make the data of this research, also for colleagues who helped and directly involved in this research, thus this research can proceed well.

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