ORIGINAL RESEARCH



Fabrication of nanosized $SO_4^{2-}/Co-AI$ mixed oxide via solution combustion method used in esterification reaction: effect of urea-nitrate ratio on the properties and performance

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Abstract

Recently, mixed metal aluminate as a highly active, stable, and reusable nanocatalyst was extremely used in chemical reactions such as esterification. The preparation of the samples via a simple, rapid, and cost-effective method is one of the main problems on the front of the industrial application of these materials. For this purpose, the solution combustion method (SCM) was assessed for the fabrication of spinel $CoAl_2O_4$, and the effects of fuel-to-oxidizer (F/O) ratio on crystalline structure, surface functional groups, surface area, morphology, and porosity were analyzed. The activity of the samples, which was impregnated by the sulfate group was evaluated in the esterification reaction. Morphology and texture properties of the samples indicated the crystallinity, surface area, and pore diameter of $CoAl_2O_4$ were enhanced by increasing the F/O ratio. It can be related to increasing the combustion reaction time and temperature. However, at F/O ratios above two, due to limited diffusion of oxygen into the reaction medium, the reaction terminated immaturely. Sulfated cobalt aluminate fabricated at F/O ratio of two was successfully synthesized in nanoscale with same particle size. Moreover, it did not show impurity in its structure that is related to the ability of the combustion method for the fabrication of pure nanocatalyst. Evaluation of the catalytic activity of the SO₄²⁻/CoAl₂O₄ nanocatalysts in the esterification reaction confirmed the results of physicochemical studies. For instance, the sample prepared at F/O ratio of two presented the highest conversion (96.7%) at methanol/oleic acid molar ratio of nine, catalyst amount of 3 wt. %, and reaction temperature and time of 120 °C and 4 h, respectively. In addition to preserving its activity for four runs (conversion > 85%), its activity was simply recovered upon regeneration by sulfate group, making it a promising alternative for industrial production of biodiesel.

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Graphic abstract



Effect of Fuel-to-Oxidizer Ratio on the Structural and Performance of Spinel CoAl₂O₄ Promoted by Sulfate Groups Used in the Esterification Reaction

Keywords Cobalt aluminate \cdot Nanocatalyst \cdot Solution combustion method (SCM) \cdot Fuel to oxidizer ratio \cdot Esterification \cdot Biodiesel

Introduction

Upon increased attention to air pollution and climate changes resulted from hydrocarbon emissions, governors are trying to provide laws for lowering the consumption of petroleum fuels by replacing them with renewable fuels [1]. A green fuel with similar properties to diesel fuel is hugely desired because of producing lower levels of hydrocarbon emissions and reducing the problem of compatibility the green fuel with the diesel engine. Biodiesel (fatty acid alkyl esters) is produced either upon esterification of free fatty acids (FFAs) using acid catalysts or transesterification of triglycerides with base catalysts. Although the transesterification reaction is the general process for conversion of biomass to biodiesel, the sensitivity of this process to the quality of the oil is a limitation. Due to remarkably increasing the biodiesel production cost using crude vegetable oils, low-cost feedstock such as inedible oils, which have high FFA contents, are suggested. However, alkali transesterification reaction cannot utilize due to soap formation by the reaction between FFA and base catalyst. Therefore, the oils contained high amounts of FFAs must react with a short-chain alcohol for reducing their FFA content for biodiesel production in which an acid catalyst is usually used. In this reaction as called esterification reaction, strong mineral acid catalysts such as H_2SO_4 , HF, and HNO_3 are generally used [2]. Although a pre-step (esterification) was recommended to decrease the FFA content of oils [3], disadvantages associated with homogeneous acid catalysts, such as corrosion, massive wastewater production during neutralization of final product, and salt generation, have increasingly led researchers toward cleaner technologies, e.g., heterogeneous acid catalysts [2].

Because of their particular advantages, heterogeneous acid catalysts have been significantly evaluated to replace homogeneous acid catalysts [4, 5]. Specially designed heterogeneous catalysts offer high activity and selectivity along with simple recovery from reaction medium without waste production as well as reusability [6, 7]. Alumino-silicate [8], $SO_4^{2-}/La^{3+}/C$ [9], and $ZnAl_2O_4$ [10], respectively, present the ability for using for 3, 10, and 4 times in the reaction. Many solid catalysts with the acidic property have been studied for the esterification of FFAs, such as sulfated metal oxides [11], sulfated carbon [12], ion-exchange resins [13] and acid ionic liquids [14]. Although, the solid acid catalysts have successfully converted a high amount of FFA to ester during esterification reaction,

some reaction conditions were beyond the standard range for the esterification reaction via homogeneous acid catalyst (reaction temperature of 60–120 °C, methanol/FFA molar ratio (MFR) of 6–12, catalyst concentration of 3–10 wt %, and reaction time of 2–4 h). Saravanan et al. [15, 16] used a methanol/FFA molar ratio of 20 at catalyst (sulfated zirconia) concentration of 6 wt % for 7 h of reaction time to obtain a yield of about 90%. Duan et al. [11] performed esterification reaction of octanoic acid at 160 °C using 50 wt % SO₄^{2–}/Al₂O₃–SiO as a catalyst for 3 h and ended up with a yield of 93%. Sulfated Zr-KIT-6 (20) was used for esterification of oleic acid, and a conversion yield of 96% was reached at 120 °C, 4 wt % of catalyst, 20 MFR, 6 h of reaction time [17].

However, leaching the active phases (especially for the catalysts impregnated by sulfate group) and deformation of the support structure (mixed metal oxides) after uses in the reaction are the major drawbacks of some of these catalysts.

Recently, several studies have pointed out high stability and activity of mixed metal aluminate catalysts in esterification and transesterification reactions [18, 19]. Metal iondoped aluminum provides large surface area and high thermal and mechanical stability as well as chemical resistance, making it promising support [20, 21]. Zinc aluminate and copper aluminate nanocatalysts were one of them that have been evaluated in the esterification of oleic acid [10]. Moreover, the reactions have been improved by the fabrication of a novel Zn_{1-r}Cu_rAl₂O₄ nanocatalyst, thereby achieving a yield of as high as 96.9% [22]. SO₃H–ZnAl₂O₄ solid acid catalyst was in the esterification reaction, and the highest FAME yield was achieved (94.59%) at 60 °C, 1.5 wt % of catalyst, 9 MFR [23]. Although the properties of MgO/MgAl₂O₄ [24] and KOH/calcium aluminate nanocatalysts [25] were thoroughly studied in the transesterification reaction, to the best of our knowledge, no study has been performed on the activity of cobalt aluminate $(CoAl_2O_4)$ as support in the esterification reaction.

Solution combustion method (SCM) as one of the effective methods was recently utilized to synthesize powders, extensively [26, 27]. It has been reported that SCM is an appropriate method for fabrication of nanocatalyst via a fast and straightforward synthesis approach that can be a suitable choice for reactions in the liquid medium such as biodiesel production. The nanosized catalyst can purely be synthesized via SCM with appropriate structural, textural properties, and performance [24].

In the present study, cobalt spinel (CoAl₂O₄) was synthesized by SCM, and as a critical parameter in SCM, fuel-to-oxidizer (F/O) ratio was optimized based on its influence on the properties and performance of CoAl₂O₄ in the esterification reaction. Sulfated CoAl₂O₄ nanocatalysts were characterized by X-ray diffraction analysis (XRD), thermogravimetric (TG) studies, Fourier-transform infrared spectroscopy (FTIR), Brunauer–Emmett–Teller and Barrett-Joyner-Halenda (BET–BJH) analysis, field emission scanning electron microscopy (FESEM), and Energy-dispersive X-ray spectroscopy (EDX), and actually examined in the esterification of oleic acid. Finally, the stability of the optimal sulfated cobalt aluminate was evaluated to assess its application for biodiesel production on an industrial scale.

Materials and methods

Catalyst preparation

For the preparation of $SO_4^{2-}/CoAl_2O_4$ as a nanocatalyst, firstly, cobalt spinel (CoAl_2O_4) was synthesized as support via solution combustion method. In this method, nitrate type of metal salts are dissolved in distilled water and appropriate amounts of fuel determined according to propellant chemistry, i.e., the ratio of oxidizer to reducer valences of components, was then added. Therefore, 0.02 mol of aluminum nitrate nonahydrate (Al(NO_3)_3.9H_2O, 99%, Merck) and 0.01 mol of cobalt nitrate hexahydrate (Co(NO_3)_2.6H_2O, 99%, Merck) were mixed into 40 mL of distilled water and then urea (NH₂CONH₂, 99%, Merck) as fuel was added at various stoichiometric ratios (0.5, 1, 1.5, 2 and 2.5). Based on the stoichiometric ratio of urea (6.67 mmol), the reaction equation is presented as follows:

$$Co(NO_3)_2.6H_2O + 2AI(NO_3)_3.9H_2O + 6.67Y CO(NH_2)_2 \rightarrow CoAI_2O_4 + 6.67Y CO_2 + (24+13.34Y) H_2O + (4+6.67Y) N_2, Y = 0.5, 1, 1.5, 2 and 2.5,$$
(1)

where Y is a proportion of F/O compared to the stoichiometric ratio.

After heating the mixture on a hotplate at 80 °C, excess water was evaporated, and a viscous gel was obtained. The gel was then put into an electric furnace at 350 °C. After evaporation of the residual water of the gel, a white smoke released indicating that the solution has reached the point of spontaneous combustion. Subsequently, an exothermic reaction with visible flame began at auto-ignition temperature. Finally, a low-density foamy solid powder was obtained, which was further crushed in a ceramic mortar and set in a desiccator. Based on the used F/O, the obtained substances were labeled as $CoAl_2O_4$ (F/O = 0.5), $CoAl_2O_4$ (F/O = 1.5), $CoAl_2O_4$ (F/O = 2) and $CoAl_2O_4$ (F/O = 2.5).

In the second step, sulfate groups were loaded on the synthesized supports via impregnation method. For this purpose, 10 mL of an aqueous solution of H_2SO_4 (1 M) was mixed with 1 g of each cobalt spinel (CoAl₂O₄) support and



the mixture was stirred for 3 h under reflux at 80 °C. Then, the mixture was placed in an oven at 110 °C for overnight to dry and calcined at 550 °C for 3 h in air ambient. The samples were labeled as S/CoAl₂O₄ (*F/O*=0.5), S/CoAl₂O₄ (*F/O*=1), S/CoAl₂O₄ (*F/O*=1.5), S/CoAl₂O₄ (*F/O*=2) and S/CoAl₂O₄ (*F/O*=2.5).

Catalyst characterization

The XRD was detected using Cu K_{α} radiation in the range of 10-60° using UNISANTIS/XMD 300 apparatus operating at 45 kV and 80 mA to determine the crystalline phase of the samples. Moreover, the Scherrer equation was utilized for calculating the crystalline size of the samples. The quality of decompositions of presursores during the synthesis of supports via combustion reaction was assessed by TG analysis (Evolution STA, SETARAM, France) in which the temperature was increased from ambient to 900 °C at a heating rate of 20 °C/min. The FTIR for evaluating the active surface functional groups of the nanocatalysts was performed using SHIMADZU 4300 (Japan) spectrometer in the range of 400-4000 cm^{-1} . The textural properties containing the specific surface area, mean pore size and pore volume of the sulfated nanocatalysts along with adsorption-desorption hystersis and pore size distribution plots were measured by BET-BJH method using PHS-1020 (PHSCHINA, China) apparatus. The morphology, particle size, external porosity, and surface structure of the nanocatalysts were studied using FESEM images pictured by MIRA3 FEG-SEM (TESCAN, Czech Republic) instrument. Moreover, the EDX technique using VEGA II Detector (Czech Republic, TESCAN) was utilized for assessing the distribution of elements in the samples. The acid strength of the solid nanocatalyst (H_0) was determined using Hammett indicators. Methyl red ($H_0 = 4.8$), *p*-dimethylaminoazobenzene ($H_0 = +3.3$), violet crystal $(H_0 = 0.8)$ and dicinnamalacetone $(H_0 = -3.0, \text{ basic})$ were utilized as Hammett indicators. The acid strength of samples was determined by the titration method using 0.01 N Butyl-Amin solution [28].

Esterification reaction

The activity of sulfated cobalt spinel nanocatalysts was assessed in the esterification reaction of oleic acid. Because of the high percentage of oleic acid ($C_{18}H_{34}O_2$, 9-octadecenoic acid), as an unsaturated fatty acid, in both edible and non-edible oils, it was selected for the esterification reaction. 20 g of oleic acid, 0.6 g of catalyst (3 wt %), and 25.8 mL of methanol (methanol-to-oleic acid molar ratio of 9) were poured into a 100 ml stainless steel batch reactor equipped with a thermocouple (Type K) and a magnetic stirrer. The esterification reaction was carried out at 90 ± 2 °C and 120 ± 2 °C for 4 h in which the mixture was stirred at



constant stirring rate of 600 rpm. Following each run, the reactor was cooled down to room temperature. Once the mixture was settled in a separation funnel, the top layer containing methyl ester (ME) was separated and heated to purify ME from excess methanol and water. Accordingly, the conversion was determined based on reducing the acidity of oleic acid, as measured by standard titration method using 0.1 M alcoholic potassium hydroxide and phenolphthalein, according below equation:

$$AI = 5.61 V / W.$$
 (2)

where AI is acid index, V is the volume of alcoholic potassium hydroxide consumed during titration and W is the weight of ME sample.

Results and discussion

XRD analysis

XRD plots of $CoAl_2O_4$ as support synthesized by SCM at different *F/O* ratios are illustrated in Fig. 1a. It has been reported that properties of the powder synthesized by SCM such as crystallite phase and size, textural properties, and particle shape and size are generally organized by the *F/O* ratio [29]. The samples present the spinel structure of $CoAl_2O_4$ (JCPDS No. 3–0896) that its crystallinity increases by more loading urea (i.e., increasing *F/O* ratio). Although cobalt oxide structures (Co₃O₄, JCPDS No. 80–1535) can observe, these peaks reduce and disappear by increasing the *F/O* ratio (especially more than 1.5). changes in the peaks could be related to the diffusion of cobalt ions into alumina lattice as a result of increased temperature and duration of combustion reaction [27, 30].

However, the cobalt oxide structure was observed at F/O ratio of 2.5. Indeed, rapid oxygen diffusion from the air is a requirement for complete combustion in fuel-rich condition [31]. However, the combustion reaction terminated immaturely because of limited diffusion of oxygen from the air into the reaction medium as a result the huge volume of smoke released from the reaction medium [32]. Therefore, the diffusion of Co ions into alumina lattice was not occurred.

Figure 1b shows XRD patterns of the sulfated cobalt aluminate. The cobalt sulfate ($CoSO_4$, JCPDS No. 72–1455) and aluminate sulfate ($Al_2(SO_4)_3$, JCPDS No. 80–1535) structures formed on the cobalt aluminate. Well-bonding of sulfate groups with the surface of support fabricated at higher *F/O* ratio (over 1.5) can be related to higher



Fig. 1 XRD patterns of a $CoAl_2O_4$ and b sulfated $CoAl_2O_4$ prepared via solution combustion method at different F/O

Nanocatalyst	BET (m ² /g)	$P_V (cm^3)$	P _d (nm)	Acidity strength		Relative crystallinity ^a		Crystalline size (nm) ^b	
				H_0	(mmol/g)	CoAl ₂ O ₄	S/CoAl ₂ O ₄	CoAl ₂ O ₄	S/CoAl ₂ O ₄
$CoAl_2O_4 (F/O = 0.5)$	-	_	_	_	_	34.9	-	-	_
$\operatorname{CoAl}_2\operatorname{O}_4(F/O=1)$	-	-	-	-	-	100	_	25.2	-
$CoAl_2O_4 (F/O = 1.5)$	-	-	-	-	-	88.4	_	24.8	-
$\operatorname{CoAl}_2\operatorname{O}_4(F/O=2)$	-	-	-	-	-	67.4	-	23.7	_
$CoAl_2O_4 (F/O = 2.5)$	-	-	-	-	-	58.2	_	20.4	-
$S/CoAl_2O_4 (F/O = 0.5)$	17.85	0.076	3.3	3.3-0.8	0.323	-	30.3	-	_
$S/CoAl_2O_4 (F/O=1)$	32.60	0.125	4.7	3.3-0.8	0.293	-	100	-	31.6
$S/CoAl_2O_4 (F/O = 1.5)$	40.96	0.348	5.6	3.3-0.8	0.340	-	83.3	-	29.3
$S/CoAl_2O_4(F/O=2)$	32.85	0.102	7.4	3.3-0.8	0.338	-	75.8	-	28.5
$S/CoAl_2O_4 (F/O = 2.5)$	29.71	0.101	6.5	3.3-0.8	0.329	-	81.8	-	25.2

Table 1	Physicochemical	properties of th	ne samples prepared	via solution	combustion metho	od at different F/O
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^aRelative crystallinity: XRD relative peak intensity at $2\theta = 36.8^{\circ}$ for $CoAl_2O_4$ and 25.1° for S/CoAl_2O_4

^bCrystalline size estimated by Scherre's equation at $2\theta = 36.8^{\circ}$ for CoAl₂O₄ and 25.1° for S/CoAl₂O₄

crystallinity of spinel CoAl₂O₄. Moreover, S/CoAl₂O₄ (F/O = 0.5) and S/CoAl₂O₄ (F/O = 1) exhibit transformed structures from amorphous to crystalline form by calcination at 550 °C.

Physicochemical properties of all samples (spinel $CoAl_2O_4$ and its sulfated structure) are tabulated in Table 1. The relative crystallinity decreases by increasing the amount of fuel beyond the stoichiometric ratio

because of the reduction in the growth of crystals of the samples. Crystalline size of the samples approves the described results, implying that $S/CoAl_2O_4$ (F/O = 1.5) and $S/CoAl_2O_4$ (F/O = 2) exhibit small crystalline size along with single-phase structure ($CoAl_2O_4$).

TG analysis

Plots of TG analysis on $CoAl_2O_4$ samples fabricated at various *F/O* ratios are presented in Fig. 2. It is well known that results of TG analysis can be used to detect unreacted raw material during a combustion reaction. For this research, the entire sample weight loss range was subdivided into three



Fig. 2 TG plots of sulfated $CoAl_2O_4$ prepared via solution combustion method at different *F/O*

Fig. 3 FTIR spectra of sulfated $CoAl_2O_4$ prepared via solution combustion method at different *F/O*

intervals, namely 50–200 °C, 200–500 °C, and 500–900 °C. The intervals can respectively refer to evaporation of physically adsorbed water on the surface of support [33], pyrolysis of organic groups and/or nitrate precursors [34], and incorporation of cobalt ions in the host material lattice to form $CoAl_2O_4$ or transition of cobalt and alumina from amorphous to crystalline phase [35].

Among other samples, CoAl_2O_4 (*F*/*O*=0.5) showed the highest weight loss in all of the regions due to less extent of decomposition of nitrate precursors to crystal form during the combustion reaction [31]. On the other hand, less weight loss can observe for the samples fabricated at higher *F*/*O* ratio because of higher combustion reaction temperature and duration [33]. The results were in agreement with those of XRD analysis, implying that a higher amount of urea must be used to have the combustion reaction completed. Among others, CoAl_2O_4 (*F*/*O*=2) shows the smallest weight loss, possibly because of the adoption of the optimum amount of fuel for preparing spinel CoAl₂O₄ as support.

FTIR analysis

The FTIR spectra of S/CoAl₂O₄ nanocatalysts are depicted in Fig. 3. Plots of all samples exhibit a broad band after 3000 cm⁻¹ that is related to O–H vibration of moisture adsorbed on the surface of catalyst as well as a peak at 1640 cm⁻¹ [22]. The peak observes at 3550 cm⁻¹ for the samples S/CoAl₂O₄ (*F*/*O*=0.5), and S/CoAl₂O₄ (*F*/*O*=1) corresponds to OH groups in unreacted raw materials [36]. Al-O bands related to stretching vibrations of AlO₆ and AlO₄



structures detect in 500-700 cm⁻¹ and 700-850 cm⁻¹ bands, respectively [33]. The Co–O vibration peak observes below 500 cm⁻¹ [30, 31]. The vibration peaks related to sulfate groups containing S=O and S–O observe in the range of 1000-1400 cm⁻¹ [11, 37].

A broad band of sulfate groups observed in S/CoAl₂O₄ (F/O = 0.5) and S/CoAl₂O₄ (F/O = 1) spectra becomes narrower by increasing the F/O ratio because of increased availability of metal ions for bonding with sulfate groups. The broad bands could be attributed to the presence of remarkable amounts of free sulfate ions with T_d symmetry on the catalyst surface, resulting in less interaction between sulfate groups and metal ions [15, 38]. On the other hands, symmetric and asymmetric stretching vibrations of S–O band for the samples fabricated at high F/O ratio could detect at 995 cm⁻¹ and 1070 cm⁻¹, respectively [39, 40]. The position of sulfate groups peaks can confirm the chelating bidentate form of bonding between SO₄^{2–} and Co-Al oxide surfaces [21, 41].

BET–BJH and acidity analyses

BET results, as listed in Table 1, showed that S/COAl₂O₄ (F/O = 1.5) has the largest surface area, followed by S/ COAl₂O₄ (F/O = 2). The BET surface area of the samples increased from 17.85 cm⁻¹ at F/O ratio of 0.5–40.96 cm⁻¹ at F/O ratio of 1.5. The increase could be due to the nourmous amount of exhausted gas and the long combustion time during catalyst preparation [25]. However, a further increase in F/O ratio caused an excessive increase in reaction temperature and time, thereby removing crystals from the support lattice, as seen in XRD analysis results [31]. Although large surface area contributes to higher catalytic activity remarkably but pore diameter plays a significant role in reactions involving reactants with large molecules. In the biodiesel production process, the effect of porosity of catalyst is even more extensive than that of surface area it was reported that pore diameter must be larger than 6 nm to facilitate the permeation of triglycerides macromolecules inside the pores to access available surface area [42]. Among other samples, S/ $COAl_2O_4$ (*F*/*O* = 2) nanocatalysts showed the highest average pore diameter (7.4 nm).

Adsorption–desorption hysteresis curves and pore size distributions of S/CoAl₂O₄ nanocatalysts prepared at various F/O ratios are depicted in Fig. 4a. Based on the figure, the samples, except S/COAl₂O₄ (F/O = 0.5) and S/COAl₂O₄ (F/O = 1.5), show characteristic curves of type IV material, as per IUPAC classification, which corresponds to mesoporous materials. In addition, their hysteresis cycles approximately are type H3, i.e., related to the plate-like particles that providing slit-shaped pores [10]. The samples prepared at F/O ratio of 0.5 and 1.5 present Type III related to nonporous material in which the pore volume and average pore size data can be referred to voids between the particles. This property can immensely effect on the catalytic activity.

In addition, the pore size distribution of the samples (Fig. 4b) indicates that pore diameter increases with increasing the F/O ratio due to the release of huge volumes of smoke during CoAl₂O₄ preparation. In S/CoAl₂O₄ (F/O=2) nanocatalyst, the pores in the range of 6–14 nm have a high volume which facilitate the motion of reactant molecules through the porosities.

Based on Table 1, the sample prepared at high *F/O* ratio (higher than stoichiometric ratio) exhibited higher acidity due to higher crystallinity of the support, thereby contributing to the formation of sulfate-metal ions such as $CoSO_4$ and $Al_2(SO_4)_3$ structures. It is well known that homogeneous mineral acid catalyst such H_2SO_4 is usually used in the esterification reaction. In the esterification reaction, protonation of the carbonyl oxygen by acid makes the carbonyl carbon a much better electrophile and it participates in the making complex with the OH group of an alcohol. Finally, water eliminates and leads to the protonated ester, and the ester is then deprotonated. Note that the acid serves two purposes.



Fig. 4 a N_2 adsorption-desorption hysteresis and b pore size distribution plots of sulfated $CoAl_2O_4$ prepared via solution combustion method at different *F/O*



First, it makes the carbonyl carbon a better electrophile and also allows for the loss of H_2O as a leaving group. Therefore, the acidity content of the catalyst is important and directly effects on the activity of a catalyst in the esterification reaction [43].

It must be attended that, although the catalyst with higher acidity presents higher activity, many factors including surface area, pore volume, and crystallinity tends to affect sample activity.

FESEM analysis

FESEM images of $S/CoAl_2O_4$ nanocatalysts prepared by SCM at various F/O ratios are illustrated in Fig. 5. The sample fabricated at F/O ratio of 0.5 exhibits an amorphous and agglomerated structure, which is attributed to insufficient

combustion reaction temperature and time. By loading more urea as fuel (F/O = 1), well-shaped particles were formed. However, the resultant structure was denser. Crystal and particle growth continued with increasing the F/O ratio to up to two, where optimal particle arrangement and diameter were observed. However, higher F/O ratio (i.e., fuel contents) suppressed the diffusion of oxygen from the air to combustion reaction medium, ending up with incomplete reactions.

EDX analysis

Results of EDX analysis of S/CoAl₂O₄ (F/O = 2) nanocatalyst are shown in Fig. 6a. Accordingly, the sample contains Al, Co, S, and O elements, with no impurity. The S/ CoAl₂O₄ (F/O = 2) nanocatalyst shows similar distribution percentages of the elements of Al, Co, and S to those of the



Fig. 5 FESEM images of sulfated $CoAl_2O_4$ nanocatalysts prepared via solution combustion method at different *F/O*



Fig. 6 a EDX and b TEM images of $S/CoAl_2O_4$ (F/O=2) nanocatalyst prepared via solution combustion method

parent solution (15.4%, 16.8%, and 16.2% for Al, Co, and S, respectively). It can confirm well fabrication of nanocatalyst without any loss of nitrate precursors during the combustion reaction.

The TEM image of the nanocatalyst is also presented in Fig. 6b. The particles show a spherical shape with narrower particle size distribution.

The catalytic activity of samples

The activity of the samples in the esterification reaction at MFR of 9 and catalyst amount of 3 wt % was evaluated at two reaction temperatures (90 °C and 120 °C) for the duration of 4 h. The results are presented in Fig. 7. It observes that all of the samples can convert a high amount of oleic acid to its methyl ester at the reaction temperature of 120 °C. It implies high activity of the CoAl₂O₄ as support in the esterification reaction, as indicated by superior conversion vields compared to results of similar studies under the same set of conditions [11, 39]. However, to better evaluate the catalytic activity of the samples, the esterification reaction was carried out at a lower reaction temperature (90 °C). As expected, due to the lower reaction rate, lower conversion was obtained. The results reveal that $S/CoAl_2O_4$ (F/O=2) nanocatalyst exhibits the highest activity, possibly because of their higher degree of crystallinity, larger pore diameter and surface area, and higher acidity.

Reusability of S/CoAl₂O₄ (F/O = 2) nanocatalyst

The reusability of S/CoAl₂O₄ (F/O=2) as the optimal nanocatalyst was examined by frequently using the samples in successive runs of esterification reaction. After each run, S/CoAl₂O₄ (F/O=2) was washed twice with methanol to eliminate unreacted reactants and products from its pores. Then, the sample was dried at 110 °C for 12 h and calcined at 550 °C for 1 h to ensure discharge of porosities. As shown in Fig. 8, the results indicate a reduction the activity of the sample in the second run. It can be related to the gradual removal of sulfate groups with weak bonds from the sample surface as a result of leaching. However, the loss in activity was lower than that reported in some studies on other sulfated catalysts [44]. However, in the next uses, less reduction was observed that can prove the stability of support and well-bonding of sulfate group with support cations. Although the leaching effect was adversely indispensable in this catalyst, regeneration of the catalyst by impregnation with sulfuric acid could improve the results of the reused catalyst.

The activity of the regenerated sample (R-5 in Fig. 8) increased sharply, so that conversion of 96.3% was achieved; i.e., the activity of the sample was comparable to that of the sample in the first use. These results suggest that S/CoAl₂O₄ (F/O=2) nanocatalyst can be successfully reused at least for four times and simply regenerated for returning its activity, making the nanocatalyst a more environmentally friendly alternative for biodiesel production.

Comparison of the results

The catalyst preparation conditions via this method, along with the activity of the optimum sample in the esterification reactio, was compared and listed in Table 2. Hashemzehi et al. [10] presented that spinel $ZnAl_2O_4$ and $CuAl_2O_4$ fabricated by combustion method had medium activity in the esterification reaction of oleic acid. On the other hands, Wang et al. reported better activity of spinel zinc aluminate by impregnation via the sulfated group. However, they







Table 2Comparison thepreparation conditions andcatalytic activity of the S/ $CoAl_2O_4$ (F/O = 2) with othercatalysts in the esterificationreaction

Catalyst	Catalyst preparation conditions				rificatio	References			
	PM	Pt	CTP (°C)	RT	CA	MFR	Rt	Y	
SO4 ²⁻ /CoAl2O4	Combustion-impregnation	5	550	120	3	9	4	96.7	This study
SO4 ²⁻ /ZnAl ₂ O ₄	Sol gel-impregnating	30	600	65	5	25	8	70	[21]
Ni/ZnAl ₂ O ₄	Hydrothermal	20	450	280	2.5	20	6	50	[45]
CuAl ₂ O ₄	Combustion	2	-	180	3	9	6	94	[10]
Aluminosilicate	Impregnation	10	120	150	3	6	9	89	[8]
SO4 ²⁻ /La ³⁺ /C	Impregnation	20	400	62	0.75	9	5	98.4	[<mark>9</mark>]
SO ₄ ²⁻ /Fe ₃ O ₄	Impregnation	30	500	105	10	1	4	68	[46]
Mg/Al	Coprecipitation	20	500	110	3	6	6	66.1	[47]

PM Preparation method, *Pt* preparation time, *PCT* calcination temperature of preparation, *RT* reaction temperature, *CA* catalyst amount, *MOR* methanol-FFA molar ratio, *Rt* reaction time, *Y* yield

used more time for fabrication the catalyst. $Ni/ZnAl_2O_4$ also showed less activity even at the reaction temperature of 280 °C [45]. The results of other researches also exhibit that the combustion method requires less time for fabrication of the final powder and the prepared nanocatalyst has high activity in the esterification reaction due to its appropriate properties.

Conclusion

Synthesis active, stable, and less-corrosive heterogeneous catalyst is recently concerned for production of biodiesel during esterification and transesterification reaction. For this purpose, $CoAl_2O_4$ spinel was firstly synthesized via SCM and impregnated by sulfate groups to prepare a catalyst for esterification reaction. Amount of urea as fuel in



the CoAl₂O₄ spinel preparation process was optimized by assessing its influence on structure and performance of the support and sulfated nanocatalyst. The results indicated that $S/CoAl_2O_4$ (F/O=2) nanocatalyst had the highest level of activity as a result of its large surface area, a high degree of crystallinity, large pore diameter, and small crystalline size. It converted 96.7% of oleic acid (as FFA) to biodiesel at middle esterification reaction conditions. Comparing the performance of the S/CoAl₂O₄ (F/O = 2) nanocatalyst to other catalysts studied in related literature confirms the high activity of the sample along with its straightforward and cost and energy effective synthesis procedure. Furthermore, it could be easily regenerated without any loss of activity, confirming enormous potentials of CoAl₂O₄ as a support for similar chemical reactions due to its mechanical and chemical resistance properties.

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Author contribution Mr. Azmoon and Mr. Heydari performed the experiments. They also had written the manuscript. Dr. Nayebzadeh and Dr. Saghatoleslami checked the manuscript English level and quality of the experiment and the results as a supervisor and advisor, respectively. They also analyzed the characterization results.

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