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Rational design of hierarchically porous sulfonic acid an d silica hybrid s with highly active sites for efficient catalytic biodiesel synthesis

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Fatt y acid methyl este r He ter ogeneou s ca t alyst Tran seste r ification *p* -toluen e su lfoni c acid Calorifi c valu e

Keywords : Ca sto r oi l

ABSTRACT

Catalysis is the vertebra of most of commercial processes, which utilizes chemical reactions to transform reagents into value added chemicals. Biodiesel synthesis from animal fats and edible vegetable oils via transesterification over homogeneou s ca t alyst s is recently take n into accoun t of unte nable by th e emer gin g bi ofuel indu stries, pa r ticularly by virtue of food *vs*. fuel counteraction, economic and environmental challenges blended with the feedstocks as well as catalytic systems, respectively. Therefore, present efforts concern with the preparation of a novel PTSA-Si catalyst and its relevance for biodiesel synthesis from non-food castor oil. It has been manifested from the experimental outcomes, the most relevant reaction parameters are, 5% PTSA-Si *(w/w)*, 65 °C reaction temperature, 1:11 O.M molar ratio and 10 h reaction time for 98.56% biodiesel yield. The PTSA-Si was appropriately analyzed using FT-IR, SEM, XRD, BET, TGA-DTA and TPD-NH₃ analysis. Since, castor oil and castor biodiesel were analyzed using FT-IR, ¹H &¹³C NMR analysis. Besides, biodiesel physico-chemical properties were pr edicted an d associated with ASTM fuel standards.

1 . Introduction

al design of hierarchically porous sulfonic acid and silica hybrid
active sites for efficient catalytic biodiesel synthesis
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Changement Channel State St Th e wind , sola r an d bi omass energy ar e pote ntial su bst itute in th e direction of petroleum based fuels and are excellent commercial source s of rene wable energy [1]. Th e insi stenc e of bi ofuel s as a gree n fuel is accelerated advancement in the last decennium with a view to reduce ou r obse ssion on petr oleum base d fuel s as a leadin g source of tran sport ation fuel as pe r th e Kyot o pr otoco l on emission co ntrol of gree nhous e gase s [2]. Bi ofuel s ar e co mpounds orig inate d from bi omass that could directly be a substitute for, or be used as mixture with conventional transportation fuels [3]. The biodiesel is a class of biofuel that coul d be pr epare d from variet y of feedstocks , assorted into thre e main groups (i) vegetable oils, (ii) animal fats and (iii) waste frying oils [4]. A fourth group of developing attention are the algal oils offering great oil pr odu ctiveness . Ho wever , th e ut ilization of alga l oi l as an alte rnative is stil l very finite du e to high attributed expenses [5]. Presently, cookin g oils ar e bein g used in more than 95 % of th e global biodiese l prep aration [[6](#page-9-5)]. Th e majo r sens e behind ut ilization of cookin g oi l fo r biodiese l prep aration is that they ar e ea sil y acce ssibl e from ma ssive agricu ltura l formulation enterprises. Nonetheless, this is not a favourable direction du e to th e si gni ficant pe rcu ssion s on food chai n invi tin g to th e so known as food *vs*. fuel conflict [7]. To date, considerable non food oils

have also been examined as ra w material s fo r biodiese l prep aration alon g with seed oils of ja troph a cu rcas, calo -phyllu m in ophyllum, to - bacco, rubber, mahua, karanja, castor and schleichera oleosa [[8\]](#page-9-7). Among all the oil crops, the castor oil has the considerable oil yield capa city. It offers th e higher yiel d of seed s an d oi l co mposition s in it s seeds. Th e majo r co mposition of ricinoleic acid is responsibl e fo r th e higher viscosity and density of castor oil. It has a potential to get dissolved in alcohol, whic h favour s th e tran seste r ification reaction . Th e main sense behind selection of castor oil as feedstocks for biodiesel preparation in present study is its higher production in India, lower eco-nomical cost and relevant fuel properties [[9\]](#page-9-8). The physicochemical properties of castor oil are listed in [Tabl](#page-1-0)e 1.

An inte res t of biodiese l as a gree n option to co nve ntional fuel co m prise its engine affinity and higher engine wear, high transportability, ea sil y avai lability, ec o -friendliness , be tte r co mbu stion pe rfo rmance, better energy generation, low sulfur and aromatic compositions, leadin g cetane inde x an d greate r biodegra dabilit y [[12](#page-9-9)]. Th e co mbu stion of biodiese l offers more than 90 % decrease in th e tota l unspen t hydr oca r bons an d a 75 –90 % weaken s cyclic ar omati c hydr oca rbons emission re lated to the utilization of petroleum based fuels. Biodiesel also demonstrate s a notewo rth y shrinkag e in pa rti c ulate s ma tters as well as CO with respect to fossil fuel [\[13](#page-9-10)]. The biodiesel is being prepared by using

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Tabl e 1

Physicochemical properties of castor oil [[10](#page-9-11),[11](#page-9-12)].

Sr. No.	Properties	Value
1.	FFA (%)	0.264
2.	Density (kg/m^3)	962.8
3.	Fire point $(^{\circ}C)$	335
4.	Flash point (°C)	298
5.	Cloud point $(^{\circ}C)$	15.8
6.	Calorific value (kJ/kg)	35684.5
7.	Kinematic viscosity (mm^2/s)	43.1
8.	Specific gravity	0.962
9.	Cetane number	40
10.	Iodine value $(gI_2/100 g)$	100

[C](#page-10-11)han plus (70

Chan plus (18)

Channel and Chan an homogeneous alkali and acidic catalysts like, KOH, NaOH, $\rm H_2SO_4$, HCl and $HNO₃$ [[14\]](#page-9-13). However, the biodiesel preparation using heterogeneous catalysts contributes a potential substitutive way for their eco-nomical production [\[15](#page-9-14)]. Solid acids offer several intrinsic advantages over liquid counterparts such as, easy separation, simple post treatment , less wast e emission an d higher therma l st abi lit y [\[16](#page-9-15) [,17](#page-10-0)]. Many soli d acid s have been deve loped to co mpe nsate th e co nve ntional li qui d acids catalysts. However, most of the solid acid catalysts are less stable, less active and more expensive than the liquid acid catalysts [\[18](#page-10-1)]. Solid acid catalysts offer economic and process benefits, especially for low cost feedstocks with high FFA content, enabling single step esterifica-tion and transesterification of bio oils [\[19](#page-10-2)]. A wide range of solid acids includin g he teropol yacid s [20 –22], su lfonate d meta l oxides [23 ,24], ca rbons [[25\]](#page-10-6), an d ze olite s [26] have been employed fo r biodiese l prep aration , ne verth eless , nove l material s with ta ilore d solvothe rma l stability and recyclability are still sought. Mesoporous solid acid catalysts are attractive in heterogeneous catalysis with a view to their tunable pore architectures, high surface areas, associated improved masstranspor t an d pote ntial to ge nerat e highly di spersed an d active site s [[27\]](#page-10-8). Ther e ar e se veral report s avai lable fo r th e si ngl e step biodiese l prep aration from wast e oils usin g soli d acids, includin g ca rbo n base d solid acid [28], active clay [29], Zr-SBA-15 [30] and sulfated tin oxide [[31\]](#page-10-12). The waste oils contain FFAs and triglycerides; both of them form biodiesel. Se veral be n efits of biodiese l in associ ation with petr oleum base d diesel ar e that it is enviro nment friendly an d safe r to use, as it is bein g pr epare d from su stainable resource s an d co ntain s li ttl e amount s of su lfur. Even though , th e cold flow assets , NO x di scharge an d higher pr odu ction expenses ar e majo r countenanc e that have to be defeated [[32\]](#page-10-13).

With the above considerations, a novel heterogeneous acid catalyst of silica fused p-toluene sulfonic acid was freshly prepared by chemical an d phys ica l mo d ification of toluen e an d si lic a ge l respectively . Th e sy nth esize d PTSA -Si wa s examined fo r si ngl e step biodiese l pr odu ction vi a tran seste r ification of no n food ca sto r oil. It ha s been co ncluded from th e tran seste r ification result s of no n food ca sto r oil, th e exce llent expe r imental parameters for the maximum biodiesel yield (98.56%) are, 5% PTSA-Si (w/w), 65 °C reaction temperature, 1:11 O:M molar ratio and 10 h reaction span . Th e sy nth esize d PTSA -Si ha s been assessed fo r re peat abi lit y stud y an d co mmenced to be four time su ccessfull y reusable without loss of their catalytic performance. The castor oil biodiesel has also been examined fo r thei r physic o -chemical properties estimation . It ha s been observed from th e stud y that al l physic o -chemical properties ar e foun d to be within li mit s pr escribe d by ASTM standards.

2 . Material s an d method

2. 1 . Materials

The methanol (≥99.0%, EMPLURA), silica gel (96.1%, 60–120 mesh), toluene (≥99.0%, EMPLURA), sodium sulphate (99.0%, FINAR), su lfuri c acid (99.9% , RANKEM), methylen e dichloride (99.8% , FINAR) and *n*-hexane (99.9%, FINAR) were furnished by Aashka Scientific Co., Surat, Gujarat. The castor oil was purchased from local oil trader (Malha r trading) , Surat, Gujarat, India. Al l reagen t ut ilize d in th e pr esent stud y were of an aly t ica l grad e an d employed withou t additional purifi cation .

2. 2 . Instrumental techniques

The FT-IR spectrum PTSA-Si was executed on a FT-IR spectrometer (Shimadzu) . Ho wever , th e FT -IR anal ysi s of ca sto r oi l an d ca sto r biodiesel were carried out on another FT-IR spectrometer (PerkinElmer). The ¹H NMR and ¹³C NMR analyses were done on a FT-NM R spectrom ete r (Bruke r 40 0 MH z NMR) . Th e ca sto r biodiese l (%) yield was calculated on GC analysis (YL 6500 GC, SUPELCO $\rm{C_{8}-~C_{24}}$ FAME mi xture). Th e wide angl e X -ra y di ffraction pa ttern wa s recorded on XR D (Rigaku) . Th e sp ecifi c su rface area of PTSA -Si wa s pr edicted on a porosimete r (2020, ASAP). Th e therma l strength of PTSA -Si wa s iden tified by TG A (PerkinElmer). Th e supe rficial mo rphol ogies of PTSA -Si were re cognize d by SE M (S3400 N, Hitachi) . Th e tota l bronsted acid sites in the PTSA-Si were predicted by TPD-NH_3 analysis (Chemisorb 2750).

2. 3 . Experimental

2.3. 1 . Synthesis of PTSA -Si hybrids

Th e PTSA -Si ha s been pr epare d by diminutive mo d ification of th e reported method [[33\]](#page-10-14). A 10 0 mL of toluen e an d 20 mL of co nce ntrated su lfuri c acid were tran sferred in a 25 0 mL glas s reacto r (FBF , adapte d with a dean stark assembly). The flask was heated with reflux at 160 °C for 6 h. The water droplets began to produce at the same time as a startin g of th e su lfonation reaction . Th e quantity of wate r co llected in th e dean stark collector was measured at regular interval of times. On the co mpl etion of 5 h of su lfonation , a 12.1 mL of wate r wa s me asure d an d th e su lfonation wa s extended as fa r as no fu rther wate r wa s pr oduced. On the completion of sulfonation, the heating was interrupted and reaction mass wa s acqu iesce d to cool at ambien t te mpe r ature . Th e crysta l lization of *p*-toluene sulfonic acid (PTSA) has been carried out through addition of 30 mL of di stilled water. Th e PTSA ha s been is olate d from th e reaction mass an d washed with agai n 10 mL of water. Th e PTSA ha s been purified by dissolving in small amount of hot water containing activate d charcoal . Finally, th e PTSA ha s been reco vered from th e aque - ous slurry and cooled with ice. A treatment of gaseous [HC](http://www.prepchem.com/synthesis-of-hydrogen-chloride/)lto this supper cooled solution leads to the precipitation of white crystals of PTSA (40.18 g) . Then after, an equimola r quantity of si lic a ge l wa s mixe d to thes e whit e crystals an d mechan icall y stirre d at room te mpe r ature fo r 1 h to su pport PTSA -Si vi a indu ction of inte rmutual hydr oge n bondin g [\[34](#page-10-15)]. Ultimately , a 53.2 5 g freshl y drie d ligh t gray fine po wde r of PTSA-Si was achieved. The reaction strategy for production of PTSA-Si is embossed in [Fig.](#page-2-0) 1 .

2.3. 2 . Catalyti c performanc e test

Th e tran seste r ification of ca sto r oi l wa s functioned ou t in 25 0 mL flat bo tto m flask, whic h wa s adapte d with reflux co ndenser , ma gneti c stirre r an d heatin g plat e sy stem. Th e required amount s of ca sto r oil, % PTSA-Si (*w/w*) and methanol were calculated stoichiometrically. The desired amount of castor oil was added to flat bottom flask followed by char gin g of PSTA -Si -methanol mi xture . Th e reflux te mpe r ature (6 5 °C) wa s maintained an d reaction asse mbl y were left uninte rrupted fo r re quired time period . Afte r th e desire d length of reaction time , th e reac tion mixture was carefully shifted to the partition funnel for the partition of th e produc t an d by product. Th e se p ara tin g fu nne l wa s agai n left uninte rrupted till th e co mplet e se p aration of biodiese l an d glycerol la y ers as depicted in Fig. S1. The glycerol collected at the bottom was separate d first. Th e ho t di stilled wate r an d anhydrou s sodium su lphat e were used fo r th e refi nin g of biodiese l layer. Co nsciously to inve stigate th e significance of reaction parameters on the (%) yield of transesterifica-

Fig. 1. Reaction scheme for the preparation of PTSA-Si catalyst.

tion products, the reactions were carried out by changing catalyst concentration (1, 2, 3, 4, 5 and 6% *w*/*w*), O:M molar ratio (1:9, 1:10, 1:11) and 1:12) and reaction time (6, 7, 8, 9 and 10 h) at reflux temperature (6 5 °C). Al l expe r iment s have been ex ecute d in triplicate s with a view to find deviations in the biodiesel yield (%) after each successful experiment . Th e scheme fo r th e tran seste r ification of ca sto r oi l ha s been demo nstrate d in Fig. 2 .

2.3. 3 . Biodiese l (%) yiel d estimation

The castor biodiesel (%) yields and their fatty acid methyl esters distribution were measured though GC investigation. The C_{8} – C_{24} composition biodiese l mi xture (S UPELCO) wa s involved as a me asure fo r qual itative an d quantitative pe rsi stenc e of biodiese l co mposition on GC (specification: $(30 \text{ m} \times 0.320 \text{ mm} \times 0.25 \text{ \mu m}$, FID, column: Agilent DB-Wax-123-7032)). The peak areas have been accounted for the proje ction of fatt y acid methyl esters co mposition of each ca sto r biodiese l sa mpl e afte r fatt y acid methyl esters of th e ca sto r biodiese l sa mples were identified. The conversion and (%) biodiesel yield were calculated by following formula (i) and (ii) [35].

sent the total peaks area,
$$
A_{IS}
$$
 refers to the methyl heptadecanote (inter-
nal standard) peak area, C_{IS} and V_{IS} stands for the concentration (mg/
mL) and volume of internal standard (mL) respectively. The *m* stands
for the quantity of the castor biological (mg).

Where, C stand for the fatty acid methyl ester content, Σ A repre-

Castor biodiesel yield
$$
(\%) = \frac{M_{\text{Biodissel}} \times C}{M_{\text{Oil}}} \times 100
$$
 (2)

Where $M_{Biodiesel}$ represent the quantity of refined fatty acid methyl esters obtained, M_{Oil} stand for the quantity of castor oil selected and C stan d fo r th e fatt y acid methyl este r co ntent estimate d as me ntioned through equation [\(i](#page-2-2)). It has been found from the gas chromatogram of ca sto r biodiesel, th e pr epare d biodiese l is offers se veral meth ylate d fatt y acid s like methyl ricinoleate, methyl oleate , methyl linoleat e an d methyl stearate . Th e GC -FI D chromatogram of ca sto r biodiese l is me n tioned in Fig. S2 .

2.3. 4 . Mechanis m of transesterificatio n of castor oi l over PTSA -Si

It ha s been alread y reported th e tran seste r ification of trigly cerides (oil) are being catalyzed by either acidic or alkaline catalysts. It involves thre e su cce ssive reversible reactions. It ha s been observed that in th e tran seste r ification reaction chain, th e trigly cerides is tran sformed stepwise to diacylglycerol, monoacylglycerol and followed by glycer-

(1)

Fig. 2. Reaction scheme for the transesterification of castor oil to biodiesel.

ine, whic h is associated with th e pr odu ction of 1 mo l of biodiese l at each stage [[20](#page-10-3)]. The mechanism of castor oil transesterification also follows the same sequence. These steps composed of (i) Protonation (H⁺) of on e of th e ca rbony l grou p of ca sto r oi l throug h th e PTSA -Si (ii) Nu cl eophili c incu rsion of th e methanol poin tin g to th e ca rbony l grou p that yiel d a tetr ahedral transitional produc t an d (iii) Pr oto n di splac ement (H +) an d di sco nne ction of th e tetr ahedral inte rmediat e yields biodiese l an d di acy lglycerol . Th e di splaced pr oto n repeatedly co nsume d throug h the leading PTSA-Si with a view to reproduce it for the next repetitive use. This co nso l idate d co ntinuit y will repeat by tw o time s till thorough sy nth esi s of mi xture of biodiese l an d glycerol . Th e mech anism scheme for the transesterification of castor oil is given in [Fig.](#page-3-0) 3.

3 . Result s an d discussion

Ge nerally , th e tran seste r ification of oi l demand s a suitable ca t alyst with a view to obtain desired production rates. The constitution of the ca t alyst is remain s esse ntial sinc e it re g ulate th e co nfi g uration s restri c tion s that th e ra w mate ria l must co ntain . Besides, th e reaction parame ters an d post is olation step s ar e pr e -planne d by th e attributes of th e ca t alyst employed [[36\]](#page-10-17). Ge nerally , majo rit y of oils ar e bein g tran sformed into biodiese l usin g alkali ca talyzed tran seste r ification method . Bu t ther e ar e se veral unco mmo n case s wherei n direct tran seste r ification ca nno t be ca rried out. Such case s appear in ra w ve getable oils like olive, jatropha, castor and cotton seed oil, etc. Such raw vegetable oils po ssess high free fatt y acid s (FFAs) co ntent s in thei r oi l co mpositions, as it forms soap on base catalyzed transesterification [\[10](#page-9-11)]. Therefore, it is

Fig. 3. Proposed reaction mechanism for the transesterification of castor oil to biodiesel.

a demand of time to develop an efficient heterogeneous acid catalyst to overcome such problems . Because, th e he ter ogeneou s acid ca t alyst s ca n co ncu rrently co nduct s tran seste r ification of TG s an d este r ification of FFAs to biodiesel. Hence, it is not required to isolate free fatty acids from th e trigly cerides [[36\]](#page-10-17). With th e abov e co nsi der ations, in th e pr e sent study, the biodiesel synthesis has been carried out by transesterification of non food castor oil over PTSA-Si as a solid acid catalyst. In orde r to optimize th e proces s parameters , th e biodiese l reaction s were ex amined by varying catalyst concentration $(1, 2, 3, 4, 5 \text{ and } 6\% \text{ } w/w)$, 0:M molar ratio (1:9, 1:10, 1:11 and 1:12) and reaction time (6, 7, 8, 9 and 10 h) at reflux temperature (65 \degree C). All the experiments were functioned at viscerous pressure till an attainment of the maximum conversions. Th e result s of ca sto r biodiese l yields (%) with di stinctive reaction parameters have been ta b ula rized in [Tabl](#page-4-0) e 2 .

3. 1 . Influenc e of oi l to methanol mola r rati o

neurons presentation in a contract of the state of t In pu rsuance to look into th e caus e of oi l to methanol (O:M) mola r rati o on biodiese l (%) yields , al l reaction s were ex ecute d ou t at divers e O:M molar ratio (1:9, 1:10, 1:11 and 1:12). In connection with [Tables](#page-4-0) 2 an d it ha s been co nfirmed that , th e biodiese l yiel d (%) is pr ecisely infl u ence by diversifying methanol to oil (M:O) molar ratio. The biodiesel (%) yiel d is grad ually increase d with th e increa sin g M: O mola r ratio. With 1:9, 1:10, 1:11 and 1:12 O:M molar ratio, the maximum biodiesel (%) yields were foun d to be 80.23% , 88.12% , 98.56% an d 94.23% re spectively . Th e optima l biodiese l (%) yiel d (98.56%) wa s foun d with 1:11 O: M mola r ratio. Th e backward reaction (a lcoho l ysi s of glycerol an d biodiesel) directly enhances upon keepin g O: M mola r rati o down from 1:11 . Accordingly, th e reversible nature of th e tran seste r ification is the responsible factor for the decrement in (%) yield of biodiesel at O:M molar ratio below 1:11. It has also been perceived that the reduction in the biodiesel (%) yield above 1:11 O:M molar ratio and this may in co njunction to th e pola r appa ren t of PTSA -Si an d methyl alcohol. Wher efore , it co nstantl y raises th e pola rit y of th e tran seste r ification , on increa sin g of M: O mola r ratio. In advanc e of, th e reaction mech anism of tran seste r ification ev idently acclaime d that , firs t oi l must reac t with ca t alyst . Though , as th e quantity of alcoho l increases, th e reaction mass

Tabl e 2

Result s of ca sto r biodiese l yiel d (%) with peculiar reaction parameters at 65 °C .

				Sr. No. O:M molar ratio Reaction time (h) % (w/w) PTSA-Si Biodiesel Yield ^a (%)
1.	1:9	6	5%	46.51 ± 0.93
2.	1:9	7	5%	59.36 ± 1.31
3.	1:9	8	5%	65.78 ± 0.70
4.	1:9	9	5%	72.32 ± 0.96
5.	1:9	10	5%	80.23 ± 0.91
6.	1:10	6	5%	58.98 ± 0.96
7.	1:10	7	5%	72.65 ± 0.89
8.	1:10	8	5%	79.25 ± 1.32
9.	1:10	9	5%	83.65 ± 0.78
10.	1:10	10	5%	88.12 ± 0.81
11.	1:11	6	5%	68.54 ± 0.91
12.	1:11	7	5%	80.29 ± 1.28
13.	1:11	8	5%	88.65 ± 1.21
14.	1:11	9	5%	91.32 ± 0.98
15.	1:11	10	5%	98.56 ± 0.99
16.	1:12	6	5%	66.32 ± 1.11
17.	1:12	7	5%	77.54 ± 0.78
18.	1:12	8	5%	86.32 ± 1.09
19.	1:12	9	5%	90.32 ± 1.16
20.	1:12	10	5%	94.23 ± 0.93
21.	1:11	10	1%	22.34 ± 1.31
22.	1:11	10	2%	38.35 ± 0.70
23.	1:11	10	3%	54.23 ± 1.11
24.	1:11	10	4%	82.56 ± 1.02
25.	1:11	10	6%	98.24 ± 0.98

^a (n = 3) All experiments have been carried out in triplicates.

is also starts to increase its polarity. Therefore, contrary to oil aspect, th e PTSA -Si co nstantl y di splaced to th e alcoho l aspect . Wher efore , th e connections of PTSA-Si in the direction of methanol phase become absolutely energeti c than co nne ction s of PTSA -Si in th e dire ction of oi l phase. Consequently, the shrinkage in the biodiesel (%) yield was noticed above 1:11 O:M molar ratio. The bar graph indicating an effect of O:M molar ratio approaching biodiesel (%) yield is manifested in [Fig.](#page-4-2) 4.

3. 2 . Influenc e of reaction time (h)

In pu rsuance to look into th e effect of reaction span (h) in dire ction of biodiese l (%) yield, al l expe r iment s were studie d employin g dive rsi fied span of reaction time s (6 , 7, 8, 9 an d 10 h) . In co nne ction with Tables 2 an d it ha s been pe rceived that th e 1:11 O: M mola r rati o demo nstrate s optimu m yiel d of biodiese l (98.56%) . Thus , 1:11 O: M mo lar ratio was preferred as an optimal ratio to examine biodiesel synthesi s employin g dive rsified span of reaction times. From th e obse rvations, it ca n be ev idenced that , with 6 h, 7 h, 8 h, 9 h an d 10 h reaction times, th e optima l biodiese l (%) yields observed were 68.54% , 80.29% , 88.65% , 91.32% an d 98.56% respectively . alon gside 10 h reaction time , it indicate s almost co mplet e co nve rsion of ca sto r oi l into biodiesel, thus , th e reaction time (h) wa s no t studie d beyond 10 h span . The bar graph indicating an effect of reaction time (h) approaching biodiesel (%) yield has been manifested in [Fig.](#page-5-0) 5.

3. 3 . Influenc e of PTSA -Si quantity (% w/w)

In pu rsuance to look into th e caus e of PTSA -Si quantity (% *w/ w*) on biodiese l (%) yields , al l expe r iment s were ex ecute d by shiftin g PTSA -Si concentration $(1, 2, 3, 4, 5 \text{ and } 6\% \text{ } w/w)$. In connection with [Tables](#page-4-0) 2 and it was realized that, the biodiesel (%) yield increases on increasing the catalyst quantity (% w/w). In the case of 1%, 2%, 3%, 4%, 5% and 6% catalyst quantity (w/w) , the biodiesel $(\%)$ yield were found to be 22.34% , 38.35% , 54.23% , 82.56, 98.56% an d 98.24% respectively at optimum O:M molar ratio. The PTSA-Si quantity above 5% (*w/w*) doesn' t sy mbo liz e an y noticeable improv ement in th e biodiese l (%) yiel d rather the (%) yield of biodiesel is slight decreased with 6% catalyst due to increment of the polarity of the reaction on increment of the % catalyst *(w/w)*. Accordingly, it was observed from the reaction outcomes, as catalyst quantity (% *w/w*) increases, the biodiesel (%) yield also increases. It ha s also been pe rceived that th e slight redu ction in th e biodiese l (%) yiel d with 6% ca t alyst (*w/ w*) an d this ma y in co njunction to the polar apparent of PTSA-Si and methyl alcohol. Wherefore, it con-

Fig. 4. Impact of oil to methanol molar ratio on the (%) yield of biodiesel (Reaction time : 10 h an d PTSA -Si : 5% (*w/ w*)) .

Fig. 5. Impact of reaction time (h) on the (%) yield of biodiesel (O:M molar ratio: 1:12 an d PTSA -Si : 5% (*w/ w*)) .

stantly raises the polarity of the transesterification, on increasing of PTSA -Si co nce ntr ation . In advanc e of th e reaction mech anism of tran s esterification evidently acclaimed that, first oil must react with catalyst. Though , as th e co nce ntr ation of ca t alyst increases, th e reaction mass is starts to increase its polarity. Therefore, contrary to oil aspect, the PTSA-Si constantly displaced to the alcohol phase. Wherefore, the connections of PTSA-Si in the direction of methanol phase become absolutely energetic than connections of PTSA-Si in the direction of oil phase. The bar graph indicating an effect of PTSA-Si quantity (% w/w) approaching biodiesel (%) yield is manifested Fig. 6.

3. 4 . FT-IR analysis of PTSA -Si

The FTIR spectroscopy is a precise technique being used particularly fo r identifyin g organi c chem icals in a whol e rang e of appl ications. An absorption pattern of synthesized PTSA-Si catalyst is accustomed by the existence of characteristic bands at 3439.19 cm⁻¹(stretching, intermolecular H- bonding, O–H), 3036.06 cm⁻¹ (stretching, C–H), 2918.40 cm⁻¹ (stretching, CH₃), 1629.90 cm⁻¹ (bending, C=C), 1206.33 cm⁻¹ (stretching, C–S), 1143.83 cm⁻¹ (stretching, O≡S≡O (by virtue of $-SO_3H$ group)), 1066.67 cm⁻¹ (stretching, S=0), 1016.52 cm⁻¹ (stretching, O-Si-O), 815.92 cm⁻¹ (p-substituted ben-

Fig. 6. Impact of % PTSA-Si (w/w) on the (%) yield of biodiesel (O:M molar ratio: 1:11 and Reaction time: 10 h).

zene ring) and 692.47 cm^{−1} (bending, C–H) subsequently. The FT-IR graph of PTSA-Si is mentioned in [Fig.](#page-5-2) 7.

3. 5 . Morphologica l study of PTSA -Si catalyst

An external morphology and structural orientation of PTSA-Si has been studied by scanning electron microscopic analysis. It has been observed from th e SE M micr ographs of PTSA -Si ca t alyst , th e PTSA pa rti cles ar e foun d with aggr egate s of irre g ula r size an d shapes afte r su lfona tion with su lfuri c acid . Whereas, th e si lic a ge l pa rticles ar e foun d with finer needles of pentagonal shape. It has been well reported that silica particles are found to have smoother surface. However, on treatment with p-toluene sulfonic acid, they are found with rough surface. The averag e pa rticl e size of PTSA -Si ha s been foun d in th e rang e of 1.07 –1.66 μ m . Muth u et al . (2014) have sy nth esize d su lfate d zi rconi a ca t alyst vi a so lvent free method an d eval uated fo r th e biodiese l sy nth e si s usin g neem oil. From th e SE M micr ograp h of su lfate d zi rconia, they realized that the particle size of sulfated zirconia is in the range of 20 –30 μ m an d is in an agglomerated form on ca lcination at 60 0 °C [[37\]](#page-10-18). The SEM micrographs of PTSA-Si are picturized in [Fig.](#page-6-0) 8.

3. 6 . Th e crystallin e structure of th e PTSA -Si

EXERC[T](#page-5-1)ION CONTROLLER C The texture characteristics of PTSA-Si were analyzed by X-ray diffraction analysis. The X-Ray diffractogram displayed characteristics diffraction pattern showing crystalline nature of PTSA-Si. The diffractogram displayed particular diffraction impression confirming crystalline texture of PTSA-Si. The particular peaks recognized at 2θ = 9.22 (deg.) (face-centered cubic), 2θ = 13.04 (deg.) (bodycentered cubic), $2\theta = 15.14$ (deg.) (face-centered cubic/diamond cubic), 2θ = 16.84 (deg.) (body-centered cubic/simple cubic), $2\theta = 18.10$ (deg.) (face-centered cubic/simple cubic), $2\theta = 22.12$ (deg.) (simple cubic), $2\theta = 24.08$ (deg.) (body-centered cubic), $2\theta = 24.62$ (deg.) (body-centered cubic), $2\theta = 25.60$ (deg.) (bodycentered cubic/simple cubic), $2\theta = 27.16$ (deg.) (body-centered cubic), $2\theta = 28.12$ (deg.) (face-centered cubic) and $2\theta = 76.16$ (deg.) (fac e -centered cubic) ar e associated with th e existenc e of si lic a ge l crys tals (JCPDS -29 -1129). Although , knife -edge d peak s observed at $2\theta = 31.74$ (deg.), $2\theta = 34.02$ (deg.), $2\theta = 38.42$ (deg.), $2\theta = 42.92$ (deg.) , $2\theta = 43.72$ (deg.), $2\theta = 45.80$ (deg.) and $2\theta = 52.68$ (deg.) ar e peculiarit y of *p -*toluen e su lfoni c acid (JCPDS -39 -1699). Th e wide angle X-Ray diffractogram of PTSA-Si has been demonstrated in [Fig.](#page-6-1) 9.

3. 7. Determinatio n of therma l stability

Th e therma l strength of th e PTSA -Si wa s assure d by TG A exam ina tion . Th e weight loss of PTSA -Si wa s me asure d as beha viour of rais e in te mpe r ature or heatin g rate . Th e TG A -DT A curv e of PTSA -Si di splay s a natural three stage disintegration diagram. The three steps in disintegr ation di agram ca n be pr edicted by loss of moisture , deco mposition of

Fig. 7. FT-IR spectrum of PTSA-Si.

Fig. 8 . SE M micr ographs of PTSA -Si .

Fig. 9. The wide angel X-Ray diffractogram of PTSA-Si.

sulfonic acid group (-SO₃H) and total degradation and decomposition of PTSA. The first weight loss (10.68%) is observed in temperature domain of 30 –15 0 °C is du e to di sintegr ation of absorbed moisture on ca t alyst su rface . Th e se con d weight loss (45.03%) is observed in te mpe r a ture domain of 150–400 °C is accredited with the disintegration of sulfonic acid groups (-SO₃H) of the toluene. The second derivative of DTA also co nfirm s th e di sintegr ation of su lfoni c acid groups from th e toluene. However, the third weight loss (5.49%) is observed in temperature domain of 40 0 –91 0 °C is an accoun t fo r th e deco mposition of toluene. In th e pr esent stud y al l tran seste r ification reaction s have been carried out at reflux temperature (65 °C). However, in the TGA curve of PTSA -Si , it wa s re cognize d that th e firs t weight loss (10.68%) is ob served in temperature scale of 30–150 °C is due to disintegration of absorbed moisture on PTSA-Si surface. Hence, the PTSA-Si offers a better therma l st abi lit y at reflux te mpe r ature . Fe rreir a e Sa nto s et al . (2014) have prepared a novel $\rm ZrO_2\!\!-\!\!SiO_2$ mixed oxides and evaluated for the thermal stability study. From the TGA thermogram of $\rm ZrO_2\rm-SiO_2$ mixed oxides , they foun d that th e almost co nstan t decrease of mass in th e scal e of 10 0 –20 0 °C , whic h is du e to th e deco mposition of water, impu rities, alkoxides, and unspent solvents. Although, beyond 700 °C tempe r ature , th e weight loss is accredited to th e dehydrox ilation of th e ca t alyst fram ework [\[38](#page-10-19)]. Th e TG A thermogram of PTSA -Si is depicted in [Fig.](#page-6-2) 10 .

3. 8 . Determinatio n of surfac e area of PTSA -Si

The specific surface area of PTSA-Si was executed by Brunauer-Emmet-Teller (BET) surface area analysis and the results are tabularized in [Tabl](#page-6-3) e 3 .

From th e BE T isotherm of PTSA -Si , it ha s been observed that th e ca t alyst offers a specific surface area of $9.61 \text{ m}^2/\text{g}$, pore volume of $0.99 \text{ cm}^3/\text{g}$ and pore size of 2.50 nm. The lower pore volume and pore size itself express that the existence of sulfonic acid $(-SO_3H)$ groups on

Fig. 10. TGA-DTA profile of PTSA-Si.

Tabl e 3 Results of specific surface area, pore volume and pore size of PTSA-Si.

Sr.		Catalyst Specific surface area	Pore volume $(P/P0)$	Pore size
No.		(m^2/g)	cm^3/g	(nm)
	$PTSA-Si$ 9.61		0.99	2.50

th e porous su rface of PTSA -Si ca t alyst . Th e hy steresi s of PTSA -Si ha s been foun d ty p icall y of type -IV , whic h indicate s th e presence of meso porous stru cture of PTSA -Si . This co ncl usion is also identica l with re - ported literature [\[39](#page-10-20)]. Ferreira e Santos *al*. have prepared a novel ZrO2 –SiO2 mixe d oxides an d eval uated fo r th e sp ecifi c su rface area de te rmination . They observed that th e ca t alyst offers a sp ecifi c su rface area of $1742 \text{ m}^2/\text{g}$ and BET isotherm found of type-IV indicating the presence of mesoporous structure of ZrO2–SiO2 mixed oxides [[38\]](#page-10-19). The BE T isotherm of PTSA -Si ha s been demo nstrate d in [Fig.](#page-7-0) 11 .

3. 9 . Determinatio n of acidity of PTSA -Si

The total active sites (H⁺) in PTSA-Si were measured using TPD- $\rm NH_3$ analysis and results are tabularized in [Tabl](#page-7-1)e 4.

In connection to the TPD-NH₃ spectrum of PTSA-Si, it was found that the catalyst offers a bronsted acidity of 1.49 mmol/g at 194.8 °C, whic h co nfirm s th e presence of acid /supe r acid site s on th e PTSA -Si su r face . Fe rreir a e Sa nto s et al . have pr epare d a nove l ZrO2 –SiO2 mixe d oxides and evaluated for the total acidity determination by TPD-NH_3 anal ysis. From th e TP D -NH 3 spectrum of ZrO2 –SiO2 mixe d oxides , they observed that the catalyst offers an acidity of 476 $mmol/g$, which affirmed the existence of acid/super acid sites on the sulfonated materials [[38\]](#page-10-19). The TPD-NH₃ spectrum of PTSA-Si is demonstrated in Fig. S3.

3.10 . FT-IR analysis of CO

The FT-IR spectrum of castor oil (CO) is accustomed with the presence of characteristics bands at 2926.01 cm^{-1} (CH₃ stretching), 2852.72 cm⁻¹ (CH₂ stretching), 2029.11 cm⁻¹ (C=C stretching), 1743.65 cm⁻¹ (C=O stretching), 1462.04 cm⁻¹ (CH bending), 1163.08 cm⁻¹ and 1095.57 (C–O stretching) and 725.23 cm⁻¹ (CH rocking) respectively. The FT-IR spectrum of CO is illustrated in Fig. S4 (a).

3.11 . FT-IR analysis of CB

The FT-IR spectrum of castor biodiesel (CB) is accustomed with the presence of an essential bands at 2926.01 cm^{-1} (CH₃ stretching),

Fig. 11 . BE T pr ofile of PTSA -Si .

2854.65 cm⁻¹ (CH₂ stretching), 2046.47 cm⁻¹ (C=C stretching), 1737.86 cm⁻¹ (C=O stretching), 1460.11 and 1436.97 cm⁻¹ (CH bending), 1195.87, 1172.72 cm⁻¹ and 1012.63 cm⁻¹ (C–O stretching) and 725.23 cm⁻¹ (CH rocking) respectively. The FT-IR spectrum of CB is illu strated in Fig. S4 (b).

3.12 . 1 H NMR analysis CO

The proton nuclear magnetic resonance (¹H NMR) is a spectroscopic techniqu e ge nerally used fo r stru ctura l dete rmination of organi c mo l e cules. The structure of castor oil was accustomed by an essential peaks at 5.33–5.40 ppm (unsaturated olefinic –CH=CH- protons), 4.24–4.39 ppm (CH–CH₂ protons), 4.11 ppm (O–CH₂ protons), 3.35–3.39 ppm (-OH protons), 2.50 ppm $(α$ -CH₂ protons), 2.23–2.28 ppm (β-CH₂ protons) and 2.02–2.09 ppm (CH₃ protons) respectively. The ¹H NMR spectrum of CO is illustrated in Fig. S5 (a).

3.13 . 1 H NMR analysis CB

Th e stru cture of ca sto r biodiese l wa s accu stome d by th e presence of an essential peaks at 7.28 ppm (CDCl₃ solvent), 5.28–5.50 ppm (unsaturated olefinic –CH $=$ CH- protons), 4.81–4.84 ppm (CH–CH $_2$ protons), 3.54–3.60 ppm (CH₃O-methoxy protons), 2.22–2.26 ppm (α-CH₂ protons) and 2.14–2.21 ppm (β-CH₂ protons) respectively. The ¹H NMR spectrum of CB is illu strated in Fig. S5 (b).

3.14 . 13 C NMR analysis of CO

The structure of castor oil was also accustomed by the presence of an esse ntial peak s at 171.98 –172.31 pp m (C O ca rbons), 126.58 –130.26 pp m (olefini c ca rbons), 69.7 2 pp m (C H – O ca rbons), 68.69 ppm (CH₂-O carbons), 61.73 ppm (CH₃ carbon), 36.4 5 –40.0 8 pp m (DMS O so lvent) an d 28.3 9 –36.4 5 pp m (aliphatic ca r bons) respectively . Th e 13 C NM R spectrum of CO is illu strated in Fig. S6 (a).

3.15 . 13 C NMR analysis of CB

Th e stru cture of ca sto r biodiese l is also accu stome d by th e presence of an esse ntial peak s at 173.50 –176.38 pp m (C O ca rbons), 124.27–132.90 ppm (olefinic carbons), 76.82–77.45 ppm (CDCl₃solvent), $63.28-73.62$ ppm (CH-O carbons), 51.37 ppm (O-CH₃ carbon) and 27.31–36.78 ppm (aliphatic carbons) respectively. The ¹³C NMR spectrum of synthesized CB is illustrated in Fig. S6 (b).

3.16 . Comparison of catalyti c exertion of PTSA -Si

The resemblance of catalytic activity of any catalyst is of significant important in order to ensure the activity of synthesized catalysts with existing literature [\[36](#page-10-17)]. Further, it also helps to design an economical viable production process. [Tabl](#page-8-0)e 5 represents the resemblance of catalytic acti vit y of PTSA -Si with ou tcome s of exis tin g li ter ature fo r th e equi v alent clas s of he ter ogeneou s bronsted acid ca t alyst s examined fo r biodiesel production through castor oil transesterification.

From [Tabl](#page-8-0)e 5 it was ascertained that PTSA-Si offers noteworthy catalytic ex ecution fo r th e tran seste r ification of ca sto r oi l as a mean s of biodiese l sy nth esis. In cu rrent inve stigation , th e su ggested fu ndame nta l reaction parameters foun d fo r th e ma x imu m biodiese l yiel d (%) are; 65 °C reaction te mpe r ature , 1:11 O: M mola r ratio, 5% (*w/ w*) PTSA -Si an d 10 h reaction span , fo r 98.56% biodiese l co nve rsion . It wa s also been established from [Tabl](#page-8-0)e 5, the outcomes of the current investigation are conveniently corresponding to the outcomes of existing literature for the identical class of heterogeneous bronsted catalysts, where anal ogously ri gorou s reaction co ndition s were demanded [\[42](#page-10-2) ,45 –[48\]](#page-10-21).

Tabl e 5

Comparison of catalytic activity of PTSA-Si with reported results (Feedstock: Castor oil).

3.17. Moisture absorption sensitivity of PTSA -Si

Th e sy nth esize d PTSA -Si wa s also eval uated fo r thei r moisture ab sorption se nsiti vity. Ther efore , herein , th e pr escribe d quantity of PTSA - Si wa s tran sferred in a glas s bo ttl e whic h kept in st abile we tness at room temperature for certain days with a view to permit the moisture absorption on the PTSA-Si catalyst. The PTSA-Si catalyst was weighed at re g ula r slot of times. Th e moisture absorption rate (*W*%) of th e PTSA Si catalyst was determined through following [formul](#page-8-1)a (iii).

$$
W\% = \left[\frac{56 \Delta m}{18m_o}\right] \times [100]
$$
\n(3)

Where, Δ m belongs to the increased weight and m_0 belongs to the in itial weight of th e PTSA -Si sa mple. Th e impact of moisture exposure time (h) on th e moisture absorption rate of PTSA -Si is illu strated in Fig. S7 .

From Fig. S7, it was examined that, the moisture absorption profile is co nti n uousl y increa sin g alon gside increa sin g in th e moisture re v ela tion span (h) up to 90 h. Ho wever , beyond 90 h of moisture exposure time (h), th e PTSA -Si doesn' t di splay an y notewo rth y inte nsification in it s weight . It is almost sa t urate d by th e atmo spheric moisture . Th e *p* toluen e su lfoni c acid (PTSA) su rface pr esent a on e hydroxyl (-OH) groups in conjunction to one sulfonic (-SO₃H) acid group in their framework. Therefore, due to polar composition of PTSA-Si, it holds a tendency to freely consume the moisture against steady humidity atmosphere.

3.18 . Influenc e of repetitive runs of PTSA -Si on castor biodiese l (%) yiel d

Th e pr odu ction expens e of biodiese l itself decide s thei r acce ssibl e an d larg e scal e co mme rcializ ation [36]. Th e ca t alyst reusabilit y is a co nsi derable fe ature fo r th e me asurement of an ec ono m ica l vi abi lit y of th e pr odu ction process. Ther efore , with an inte ntion to cu t down an ec ono m ica l expens e of biodiese l pr odu ction , th e PTSA -Si wa s eval uated for their possible recyclability in the castor oil transesterification. Accordingly, in pr esent work , afte r each su ccessfu l cycle, th e PTSA -Si wa s detached ou t from th e reaction batc h by mean s of va cuu m se p aration an d mediated with fe w quantity of methylen e dichloride with a view to expe l impurities like , unused ca sto r oil, glycerol an d unused alcohol. Before the repetitive use, the purified PTSA-Si was putted up in an air dryer at 100 °C for 24 h with a view to remove volatile solvent and regeneration of active centres $(H⁺)$ on the toluene surface. It was observed through execution of castor oil transesterification, the PTSA-Si coul d have a si gni ficant pote ntial to recycl e four time s left ou t charac te risti c dese rtion of ca talytic pe rfo rmance. This reusabilit y migh t be du e to th e st abi lit y of PTSA -Si an d th e strong attachment of su lfoni c acid grou p (-SO ³H) with toluene. Noneth eless , th e mino r redu ction in the (%) yield of castor biodiesel has been noticed during the repetitive cycles of PTSA-Si. An extraction of active site $(H⁺)$ or physicochemical change of PTSA -Si fram ework at 65 °C te mpe r ature coul d responsibl e fo r th e deactivation of PTSA -Si . Th e toluen e su stain s it s stru cture over the castor oil transesterification without any severe change. The virgin PTSA -Si coul d di splay s optimu m co nve rsion of ca sto r oi l to biodiese l equal to 98.56%. While, it's first, second, third and fourth repetitive run coul d di splay s optimu m tran sfo rmation s of oi l to biodiese l up to 94.25% , 90.26% , 86.28% an d 81.20% su bsequently. Li u et al . (2008) have synthesized sulfonated ordered mesoporous carbon $(\mathrm{OMC\text{-}SO}_{3}\mathrm{H})$ an d examined fo r biodiese l sy nth esi s throug h este r ification of olei c acid. From the reusability study of the $OMC-SO₃H$ catalyst, they concluded that the $\rm OMC\text{-}SO_3H$ catalyst can be reusable for four successful time s withou t noticeable loss of ca talytic exertion . They also co ncluded that the reusability is due to the thermal stability of OMC-SO₃H and the strong coupling of sulfonic acid group with carbon [[49\]](#page-10-26). An impact of PTSA-Si runs on the biodiesel (%) yield is exemplified in [Fig.](#page-8-2) 12.

3.19 . Assessment of fuel properties of castor biodiese l

The castor oil plant grows in a hot and humid environment; it has a growin g se aso n of 4 – 5 months . Th e ca sto r oi l is a colourless to very pale yellow viscous liquid with moderate or no odour or taste. Castor oil offers a very lower pour and cloud points which make castor biodiesel a good alte rnative in wi nte r se aso n [\[50](#page-10-8)]. Th e physic o -chemical proper ties of biodiesel can diversify considerably from one raw material to anothe r owin g to it s somewhat high mo l e c ula r mass than co nve ntional diesel . Th e co nsi derable fuel standard sy stems like ASTM , AOCS , IS , JIS, BS an d EU ar e frui tfu l fo r th e co rrelation of th e physic o -chemical properties of biodiesel produced from diversified oils. The physicochemical properties ar e us efu l fo r an y fuel to dete rmine thei r engine pe rfo rmance, emission , storag e an d tran sport ation s co nditions. Th e fuel

Fig. 12. Impact of PTSA-Si runs time on the (%) yield of biodiesel ((i) 1:11 oil to methanol molar ratio, (ii) 5% PTSA-Si catalyst *(w/w)*, (iii) 65 °C reaction tempe r ature an d (iv) 10 h reaction time).

properties of castor biodiesel with contrast to ASTM standards have been ta b ula rized in [Tabl](#page-9-16) e 6 .

Th e cetane nu mbe r wa s me asure d on cetane analyser (AFIDA 2805). The density of castor biodiesel was measured by hydrometer mechanism . Th e acid an d iodine va lue s have been me asure d by titr ation meth ods. Th e sp ecifi c gravit y of biodiese l ha s been ca lculate d usin g sp ecifi c gravit y bo ttles . Flas h an d fire points were pr edicted by closed cu p tester (ADITYA-SUNBIM). The refractive index has been measure by refractomete r (A nto n parr , abbema t 300) . Th e clou d an d pour points have been dete rmine d on clou d an d pour poin t an alyze r (ALA Nready, CP P 5Gs) . Th e calorifi c valu e of ca sto r biodiese l ha s been dete rmine d throug h ox yge n bomb calorimete r (Par r 6400 calorimeter) . Th e kine mati c vi sco sit y an d vi sco sit y were me asure d on vi scomete r (Adity a 01). From [Tables](#page-9-16) 6 and it was recognized that all physico-chemical properties of castor biodiesel are according to the test restrictions, which are esta blished by ASTM standards.

3.20 . Industrial aspect of th e proces s

Announce and the particular particular and the energy of the energy An economical viability is very necessary with a view to explore preparation of biodiesel at commercial spectrum [\[36](#page-10-17)[,51](#page-10-9)]. When smallscal e ma n ufa ctu rer s embark on su rve y of th e finances of biodiese l pr o du ction , they fr equentl y di scove r it co mplex to acquir e oi l at a lowe r cost for cost-effective production. The oil is the potential contributor to the cost (more than 80%) of biodiesel manufacturing [\[36](#page-10-17)[,51](#page-10-9)]. Fresh oil obtained from an oilsee d ma n ufa cture r tend s to be th e chie f quality, most co nsi stent an d ea sil y avai lable . Ho wever , it also is expected to be a very costly resource of ve getable oil. Th e biodiese l reaction absorb s 11% methanol (% w/w) in the transesterification process. Nonetheless, most ma n ufa ctu rer s ut ilize betwee n 18 an d 22 % methanol to ce rtify a more complete conversion and make use of all quantity of oil during reaction. Small-scale biodiesel production economics can vary considerably dependin g on sp ecifi c ma n ufa ctu rer's sele ction of oi l source , ma chinery, me rchandise an d th e expens e of ma npower. Ho wever , se veral fundamental controversies affect all small-scale manufacturers. Thus, Present work will deals with the usage of non-edible castor oil as an oil source fo r biodiese l prep aration , becaus e of no n -edibility, th e ca sto r oi l is cheap and readily available in the oil market. In addition, reusability of PTSA -Si will be take n in to th e account, as he ter ogeneou s clas s of ca t alyst . Th e prep aration cost fo r PTSA -Si woul d no t involv e larg e cost with a view to only single step sulfonation is carried out for its preparation . Ther efore , this proces s woul d offers an ec ono m ica l biodiese l ma n ufa ctu rin g usin g milder reaction parameters .

Tabl e 6

4 . Conclusion

In cu rrent inve stigation , a promisin g four time reusable PTSA -Si is freshly prepared and investigated for transesterification of castor oil regardin g biodiese l pr odu ction . It ha s been ma n ifested agains t th e reac tion ou tcomes; th e be tte r re l evant reaction parameters fo r biodiese l sy nth esi s as a mean s of tran seste r ification of ca sto r oi l are; 5% PTSA -Si (*w/w*), 65 °C reaction temperature, 1:11 O:M molar ratio and 10 h reaction time fo r 98.56% biodiese l yield. In view of sy nchronously co ndu c tion of an esterification and transesterification by PTSA-Si catalyst, thus , it is no t si gni ficant to se gregate FFAs from ca sto r oi l an d ther efore , it yields ec ono m ica l biodiese l with appr eciated co nve rsion .

Declaratio n of competin g interest

The authors declare that they have no known competing financial inte rests or pe rsona l relationship s that coul d have appeared to infl u ence th e work reported in this paper.

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Appendix A . Supplementar y data

Su ppl eme ntary data to this articl e ca n be foun d online at [https://](https://doi.org/10.1016/j.cinorg.2023.100005) doi.org/10.1016/j.cinorg.2023.100005 .

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Abbreviation s

y. Y. in D.E. Sometimes, [R](http://refhub.elsevier.com/S2949-7469(23)00005-8/sref47). Sometimes, R. Sometimes, D.[E](http://refhub.elsevier.com/S2949-7469(23)00005-8/sref35). Sometimes, R. Sometimes, [D](http://refhub.elsevier.com/S2949-7469(23)00005-8/sref29).E. Sometim *JCPD S :* Join t committe e on powder diffractio n standard s *OM C–SO 3 H– :* Ordere d mesoporous carbon - Sulfonic acid *PAHs :* Polycyclic aromatic hydrocarbons *HC :* Hydrocarbons *PTSA -Si : p* -toluen e sulfonic acid -silica *NMR: Nuclear magnetic resonance FFAs – :* Free fatt y acid s *SEM:* Scanning electron microscope *XR D :* X -ra y diffractio n *TGA :* DT A -Thermogravimentric analysis - Differential therma l analysis *FT-IR :* Fourie r transforme d infrared spectroscopy ASTM: American society for testing and materials *AOCS :* American oi l chemical societ y *IS :* Indian standard s *EN:* European standard s *BE T:* Brunauer - Emmett – Teller *GC -FID :* Ga s chromatography - Flam e ionization detector *JIS :* Japanese industrial standard s *BS :* Britis h standard s *CO :* Castor oi l *CB :* Castor biodiese l