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Spongy titanosilicate promotes the catalytic performance and reusability of WO₃ in oxidative cleavage of methyl oleate†

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A tungsten containing catalyst catalyzed oxidative cleavage of methyl oleate (MO) by employing H_2O_2 as an oxidant and is known as an efficient approach for preparing high value-added chemicals, however, the tungsten leaching problem remains unresolved. In this work, a binary catalyst consisting of tungsten oxide (WO₃) and spongy titanosilicate (STS) zeolite is proposed for MO oxidative cleavage. The function of STS in this catalyst is investigated. On the one hand, STS converts MO to 9,10-epoxystearate (MES), which further forms nonyl aldehyde (NA) and methyl azelaaldehydate (MAA) with the catalysis of WO₃. In this way, MO oxidation and hydrolysis that generates unwanted diol product 9,10-dihydroxystearate (MDS) decreases obviously. On the other hand, STS decomposes peroxide and promotes the conversion of soluble peroxotungstate to insoluble polytungstate. Meanwhile, these tungsten species are allowed to precipitate on its surface instead of remaining in the liquid phase owing to its relative large specific area. Therefore, tungsten leaching can be reduced from 37.0% to 1.2%. Due to the cooperation of WO₃ and STS, 94.4% MO conversion and oxidative cleavage product selectivity of 63.1% are achieved, and the WO₃-STS binary catalyst maintains excellent catalytic performance for 8 recycling reactions.

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1. Introduction

Unsaturated fatty acid esters (UFAE), such as Methyl Oleate (MO), methyl linoleate, linolenic acid methyl ester and so on, are renewable raw materials with high abundance in nature. Conversion of UFAE into aldehydes, acids or esters through oxidative cleavage of its double bond has been considered as an economic and sustainable solution to obtain high value-added chemicals, and the research has attracted considerable academic and industrial interest.¹⁻⁵

Common oxidants such as potassium permanganate and ozone can be used for oxidative cleavage reaction of UFAE. ⁶⁻⁹ However, the toxicity and potential harm to humans limit their application. Some researchers turned to use hydrogen peroxide (H_2O_2) as an oxidant for this reaction ^{10–12} as H_2O_2 is environmentally benign and produces water as the only by-product. Some transition metal containing catalysts, ^{13–16} such as H_2WO_4 @Al-MCM-41, ¹² BaFeO_{3- \hat{o}}, ^{17,18} Ru(OH)_x/ γ -Al₂O₃ ¹⁹ and VO@TiO₂ ²⁰ have been proved efficient in catalyzing this reaction with H_2O_2 as oxidant. Among them, tungsten containing catalysts such as tungsten oxide (WO₃), WO₃ nanoparticles, ²⁴

State Key Laboratory of Catalytic Materials and Reaction Engineering, Research Institute of Petroleum Processing, SINOPEC, 100083, Beijing, PR China. E-mail: pengxx.ripp@sinopec.com tungstic acid,²⁵ tungstates and supported WO₃²⁶⁻³⁰ have shown the best catalytic performance.^{4,21-23}

Although effective in catalytic oxidative cleavage of UFAE, tungsten containing catalyst suffers from active center leaching problem, and thus limits its large scale application. Take catalytic oxidative cleavage of MO by WO₃ and H₂O₂ for example, the reason for tungsten leaching has already been illuminated.31-33 WO₃ firstly reacts with H₂O₂ to form the active peroxotungstate species, which are soluble in the reaction system. Then, peroxotungstate converts MO into oxidation products and itself turns into insoluble polytungstate species. This compound could be regenerated into active peroxotungstate through reacting with H_2O_2 , so the reaction cycle can be continued. After finishing the reaction, residual H2O2 would still react with polytungstate, and part of soluble tungsten species would remain in the reaction solution which leads to the loss of tungsten and lower the reusability of the catalyst. Therefore, in order to improve the performance and reusability of WO3 catalyst, it is important to reduce the residual soluble tungsten species within the reaction system.

In this work, we report a robust binary catalyst consisting of WO_3 and spongy titanosilicate (STS) zeolite for this reaction. It is found that the addition of STS would enhance the epoxidation of MO, accelerating the consumption of H_2O_2 which promoted the precipitation of tungsten species. Additionally, the relatively large specific surface area of STS facilitates the capture of precipitated tungsten species and thus improves the

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reusability of the catalyst. Compared with WO $_3$ catalyst, WO $_3$ -STS binary catalyst demonstrated improved catalytic performance (94.4% MO conversion and >63% oxidative cleavage products selectivity) and excellent reusability (8 cycles without significant loss in catalytic performance).

2. Experimental

2.1 Materials

Methyl oleate (MO, 96 wt%) and WO₃ (diameter < 200 nm, > 99 wt%) were purchased from Alfa Aesar. Tert-butyl alcohol (t-BuOH, >99 wt%) was purchased from Tokyo Chemical Industry Co., Ltd. Tetraethyl orthosilicate (TEOS) and tetrabutyl titanate (TBOT, >98 wt%) were obtained from Changling Catalyst Co. Ltd. N-Phenyl-3-aminopropyltrimethoxysilane (PHATMS), tetrapropylammonium hydroxide (TPAOH, 25 wt%) were obtained from Tokyo Chemical Industry Co. Ltd., methyl palmitate is (MP, 99wt%), 9,10-epoxystearate (MES, 99 wt%) and 9,10-dihydroxystearate (MDS, 99 wt%) were supplied by Sigma-Aldrich. H₂O₂ (30 wt%) was purchased from Sinopharm Co. Ltd. All materials were used without further purification. Commercial TS-1 zeolite (MFI type) was supplied by Changling Catalyst Co. Ltd. Spongy silicalite-1 (SS) zeolite (MFI type) and MnO₂/ Al₂O₃(10 wt% MnO₂) was supplied by Research Institute of Petroleum Processing, SINOPEC.

2.2 Preparation of the spongy titanosilicate (STS)

STS was synthesized under hydrothermal conditions with PHATMS as the zeolite pore extending reagent according to the ref. 34. The starting materials $SiO_2: TiO_2:TPAOH: H_2-O: PHATMS$ with a molar ratio of 1:0.04:0.4:40:0.05 were heated to 170 °C for 3 days under autogenous pressure. The precipitate was collected, washed by distilled water and dried at 110 °C. After calcination at 550 °C for 6 h, the STS zeolite was obtained.

2.3 Characterization

Powder X-ray diffraction (XRD) patterns were obtained on a Philips X'pert-Pro diffractometer using nickel-filtered Cu Ka radiation. Ti/Si molar ratio was quantitatively measured by Xray fluorescence (XRF) on a Rigaku 3721E spectrometer running at 40 kV. Scanning electron microscopy (SEM) images were taken from a Hitachi4800 microscope. Transition electron microscopy (TEM) images were recorded on an aberrationcorrected microscope (JEMARM200F) working at an acceleration voltage of 200 kV. STEM EDS-mapping pictures were acquired from selected areas of the HAADF-STEM images with continuous drift correction. Nitrogen adsorption-desorption isotherms were collected at 77 K on a Quantachrome AuotosorbiQ2 apparatus. Texture properties were derived from the isotherms using Brunauer-Emmett-Teller (BET) method and tplot method. Cold-spray ionization (CSI) mass spectra were recorded on a JEOL JMS-T100CS spectrometer. Pore size distribution curves were calculated from the adsorption branch according to Brunauer-Emmett-Teller (BJH) method. Tungsten

leaching was detected by Inductively Coupled Plasma Optical Emission Spectrometer (ICP-OES).

2.4 Catalytic reaction

The reactions were carried out in a 25 mL round-bottom flask under constant temperature and pressure. The reactor was equipped with a condenser, a magnetic stirrer, and an oil bath. A typical procedure for the catalytic oxidative cleavage of MO is as follows: MO (0.5 mmol), $\rm H_2O_2$ (2.5 mmol), WO₃ (0.085 mmol), STS (10 wt% with respect to MO) and t-BuOH (20 mmol) were fed to the flask. Then, the flask was placed into the oil bath under 80 °C and reacted for 4–8 h. For the catalyst recycling reactions, the catalyst was recovered by using a centrifuge at 6000 rpm. The catalyst was directly reused in the next reaction cycle without further treatment. The products were sampled and analyzed, and the concentration of tungsten in the solution was determined by ICP-OES.

2.5 Product analysis

The consumption of H_2O_2 was determined by indirect iodine titration method. Products were determined on Agilent 9790/5875C GC-MS equipped with HP-5MS(30 m \times 0.25 mm \times 0.25 m) column. The conversion of MO and product distribution were analyzed on Agilent 9790 GC equipped with a flame ionization detector and SE-54 (30 m \times 0.25 mm \times 0.25 m) column with methyl palmitate as internal standard. The conversion and selectivity were calculated according to the following equations.

Conversion of MO =
$$\frac{n_{0,\text{MO}} - n_{\text{MO}}}{n_{0,\text{MO}}} \times 100\%$$

Selectivity of Product =
$$\frac{n_i}{n_{0,MO} - n_{MO}} \times 100\%$$

where, n_0 is the mole of compound, i refers to generated products, *e.g.* MES, nonyl aldehyde (NA), pelargonic acid (PA), methyl azelaaldehydate (MAA), methyl hydrogen azelate (MHA) and MDS.

Results and discussion

3.1 Characterization of fresh catalyst

To begin with, commercial TS-1 zeolite was used for comparison with STS zeolite and the XRD and XRF results are shown in Fig. 1. Both TS-1 and STS zeolites show typical MFI topological features in the region of 22° – 25° . Different from TS-1, STS shows no peak at $2\theta=25.40^{\circ}$, which is assigned to anatase species. The relative crystallinity of STS is lower than that of TS-1 (Table 1). It is inferred that the pore extending reagent PHATMS slowed down the MFI crystals formation and expanded the pore size in STS.³⁴ The Ti/Si molar ratio measured by XRF is very close for STS and TS-1 (Table 1).

Nitrogen adsorption-desorption isotherm as well as BET and BJH methods were employed to investigate the texture properties of TS-1 and STS (Table 1 and Fig. 2a). Both two samples show type-IV isotherms with H4 type hysteresis loops (Fig. 2a), suggesting the formation of mesopores and multi-layer

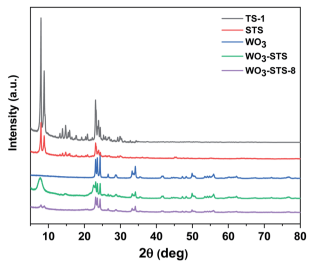


Fig. 1 XRD patterns of various catalysts

adsorption of N_2 . The relatively large hysteresis loop of STS which is mainly located at medium pressure zone (0.45 < p/p_0 < 0.9) indicates a uniform distribution of mesoporous structure. Texture properties of TS-1 and STS (Table 1) confirms the existence of mesopores. Moreover, STS shows a narrow pore size distribution at around 4 nm, which is absent from TS-1 (Fig. 2b). It could be due to the effective pore-making effect of PHATMS reagent.

SEM and TEM images of TS-1 and STS zeolites are shown in Fig. 3. Both TS-1 and STS present "raspberry-like" spherical morphology with relative smooth surface. The average particle size is about 300 to 400 nm. TEM image shows solid structure of TS-1, while STS is spongy-like morphology with intracrystalline mesoporosity owing to the aggregation of small particles. It is also found that due to the small particles of STS, the intensity of XRD is lowered down and its relative crystallinity is much lower than that of TS-1.

The XRD pattern of WO₃ displayed in Fig. 1 clearly shows monoclinic crystal phase (JCPDS no. 43-1035). Then we prepared WO₃-STS sample by mechanical mixing WO₃ and STS in a mortar with mass ratio of 15:100 and characterized it by XRD. The result in Fig. 1 shows that the peak intensity of both WO₃ and STS are decreased. Compared with WO₃, the characteristic peaks of STS is more difficult to be recognized.

3.2 Catalytic performance

SS, TS-1, STS, WO₃, WO₃-TS-1, WO₃-SS and WO₃-STS were employed as catalyst for MO oxidative cleavage reaction,

respectively, and the results are listed in Table 2. As can be seen from Table 2, there is no reaction occurred without catalyst or using SS as catalyst. When TS-1 is used, 9.8% conversion of MO is obtained and the only product is MES. This is because framework Ti species of TS-1 is effective in catalyze double bond forming epoxy bond, but its pore size is too small for the diffusion of this bulk substrate. In contrast, STS shows higher catalytic activity than TS-1 due to the better accessibility of active site, the conversion of MO reached to 30.8%, and the MES selectivity is 69.5%. It is noticed that apart from MES, some other deep oxidation products of MES like NA, PA, MAA and MHA are also observed as well as MDS, which is originated from hydrolysis of MES. WO3 is much effective for MO oxidative cleavage reaction, as MO conversion reaches to 91.8%. However, half of MES converted into MDS. Obvious WO₃ leaching is observed as 37.0% of WO3 is dissolved and lost with the liquid after reaction.

When we use mechanical mixture of WO₃ and TS-1 as catalyst, MO conversion is much higher than Ts-1 but lower than WO₃. Compared with products distribution of WO₃, the selectivity of MES decreased to 12.3%, both NA and MAA selectivity increased and the MDS selectivity also decreased slightly. It suggests that the products distribution could be mediated by combining WO₃ and TS-1. However, the WO₃ leaching is still up to 25.0% and is unacceptable for a sustainable process. The performance of WO₃-SS is basically the same with WO₃ (Table 2, entry 5 and 7) as well as the tungsten leaching rate, suggesting that SS is ineffective for the MO oxidative cleavage reaction. Further, we used the binary WO3-STS as catalyst for MO oxidative cleavage reaction. The results showed MO conversion is higher than WO₃ (91.9%). In contrast with WO3 and WO3-TS-1, the selectivity of NA and MAA is much higher and both MES and MDS decreases obviously. Among all catalysts, the highest MO conversion and total selectivity of oxidative cleavage products (NA, PA, MAA and MHA, 63.1%) are achieved by WO₃-STS. More importantly, tungsten leaching of WO₃-STS is reduced from 37.0% to 1.2%. The increased catalytic activity, enhanced oxidative cleavage products selectivity and suppressed tungsten leaching suggest synergistic effect from the combination of WO₃ and STS. The dependence of catalyst amount on the MO oxidative cleavage reaction is investigate and provided in ESI Table S1.† The results also confirms both W and Ti are essential for this reaction.

3.3 Reusability of WO₃-STS

In order to investigate the reusability of WO₃-STS, the catalyst was recovered by centrifugal separation, and reused in a subsequent reaction without drying, washing or calcination

 Table 1
 Relative crystallinity, Ti/Si ratio and texture properties of titanosilicates

Samples	Relative crystallinity (%)	Ti/Si	$S_{\rm BET}$ (m ² g ⁻¹)	$S_{\rm ext} \left({ m m}^2 \ { m g}^{-1} ight)$	$S_{ m micro} \left({ m m}^2 \ { m g}^{-1} ight)$	$V_{\rm meso}$ (cm ³ g ⁻¹)
TS-1	100	0.0355	440	48	392	0.104
STS	78	0.0350	528	129	206	0.143

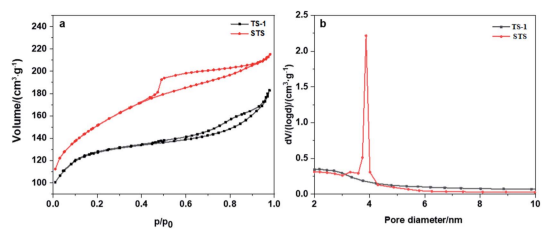


Fig. 2 (a) Nitrogen adsorption—desorption isotherms of TS-1 and STS, (b) pore size distribution of TS-1 and STS.

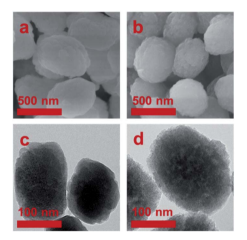


Fig. 3 $\,$ SEM images of (a) TS-1 and (b) STS, TEM images of (c) TS-1 and (d) STS.

after each reaction cycle. WO₃ was employed as control and tungsten concentration in liquid phase was measured. The results are shown in Fig. 4. Obviously, when using WO₃ as catalysis, MO conversion decreases rapidly after three cycles.

Each reaction cycle is accompanied with a serious WO₃ leaching, and thus leading to the worse reuse effect.

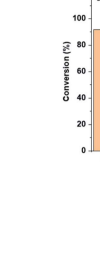
While for WO₃–STS, the story is totally different. After eight reaction cycles, the catalyst shows almost no loss of catalytic activity and oxidative cleavage products selectivity, the MO conversion even increases to 94.4% with oxidative cleavage products selectivity maintaining above 63.1%, suggesting excellent reusability of the WO₃–STS catalyst. In addition, the tungsten leaching of the WO₃–STS catalyst is controlled below 2% for each reaction cycle (Fig. 4c), and it exhibits excellent mass stability. Due to the ignorable loss of WO₃, the WO₃–STS binary catalyst can maintain excellent performance during recycling reactions.

Further, in order to study the stability of WO₃–STS catalyst, we analyzed the phase structure, morphology and micro region composition of the binary catalyst after it was recovered from the eighth reaction cycle by using XRD, STEM and EDS. The sample is denoted as WO₃–STS-8. After the eighth recycle, the recovered catalyst remained most of its original diffraction peaks as fresh WO₃–STS (Fig. 1). But the peak intensity decreases significantly especially for the peaks with diffraction angle lower than 10° which might be caused by decrease of crystal size. Nonetheless, the characteristic diffraction peaks at $2\theta = 23.07^{\circ}$, 23.54° , 24.35° , 33.18° and 34.19° which are

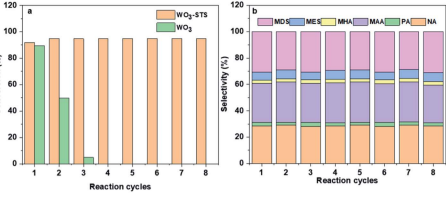
Table 2 Catalytic performance of different catalysts for the oxidative cleavage of MO^a

		Conversion (%)	Selectivity (%)						m 1 1
Entry	Catalyst		NA	PA	MAA	МНА	MES	MDS	Tungsten leaching (%)
1	_	0	0	0	0	0	0	0	_
2	SS	0	0	0	0	0	0	0	_
3	TS-1	9.8	0	0	0	0	100.0		_
4	STS	30.8	5.3	2.5	5.5	2.0	69.5	15.2	_
5	WO_3	91.8	14.2	1.1	13.1	0.9	20.2	50.5	37.0
6	WO ₃ -TS-1	76.5	19.6	0	20.9	0	12.3	47.2	25.0
7	WO_3 -SS	90.1	13.2	0.7	12.6	0.5	21.7	51.3	42.0
8	WO ₃ -STS	91.9	28.5	2.6	29.7	2.3	6.3	30.6	1.2

 $^{^{}a}$ n(MO): n(H₂O₂): n(t-BuOH) = 1: 2.5: 20, <math>m(catalyst): m(MO) = 1: 10, temperature 80 $^{\circ}$ C, reaction time 4 h.



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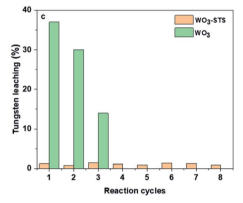


Fig. 4 Reusability of WO₃ and WO₃-STS catalysts: (a) conversion, (b)selectivity and (c)tungsten leaching. $n(MO): n(H_2O_2): n(t-BuOH) = 1:2.5:20, m(catalyst): m(MO) = 1:10, temperature 80 °C, reaction time 4 h.$

attributed to monoclinic WO_3 crystal are discernible. Although it's hard to distinguish STS from WO_3 since the characteristic peaks of STS overlap with WO_3 , it is safe to conclude that WO_3 still exists as crystal. The nitrogen adsorption–desorption isotherms and pore size distribution were obtained for fresh WO_3 –STS and the one used for 8 cycles are provided in the ESI Fig. S1.† As can be seen from the results, the isotherms and pore size distribution remain nearly unchanged. The texture properties (Table S2†) reveals that after 8 recycles, specific area of micropores and pore volume decreased slightly.

Two different morphologies are observed in the TEM image, and three areas are marked according to the TEM image contrast (Fig. 5a). The elemental mapping results reveal that O, Si and Ti atoms distribute in area 1, 2 and 3, but W mainly appears in area 1 (Table 3). The elemental distribution was further analyzed by EDS (Fig. 5b-e). It confirms that W mainly located at area 1, and Si and Ti mainly located at area 3. For area 2, all Si, Ti and W atoms are discovered since two kinds of particles is mixed. The above analysis suggests that the particles in area 1 are mainly WO₃, while the particles in area 3 are mainly STS zeolite. The intracrystalline mesopores in area 3 also proves its zeolite nature. Then, we employ HADDF characterization method to analyze the STS zeolite. It is surprise to find out that many sub-nano WO₃ particles dispersed on the STS surface with diameter of 1-3 nm (Fig. 5f-h). These particles might be generated during the MO oxidative reaction. Since the morphology of WO3 in area 1 remains well, it is inferred that only part of WO_3 involved the reaction and finally deposited on the surface of STS. This result also explains why W is discovered in area 3.

3.4 Reaction pathway

As mentioned in Table 2, WO₃ is very effective in catalyzing MO oxidative cleavage reaction, therefore, its products distribution at different reaction time was investigated and the results are displayed in Fig. 6. The MO conversion increases rapidly before 4 h and finally reaches to above 94%. The product distribution changes a lot during the reaction. At the initial stage, the main products are MES and MDS, together with a small amount of oxidative cleavage products, NA and MAA. As the reactions went on, NA and MAA increased along with the decrease of MES, and MDS also increased slightly. A small amount of PA and MHA also can be observed in the later period. The total selectivity of oxidative cleavage products is about 40% at reaction time of 8 h.

In order to figure out how the reaction occurred, we further carried out the WO₃ catalyzed experiments with epoxide MES and diol MDS as starting materials, and the results are displayed in Table 4. It can be seen that MES is quite active in this reaction, 90% of it converts into NA, MAA and MDS with selectivity 36.0%, 35.5% and 28.5%, respectively. That is to say, oxidative cleavage and hydrolysis are both possible reactions for MES but the latter is more preferred. If all MDS is generated from MES in MO oxidative cleavage reaction, then MDS

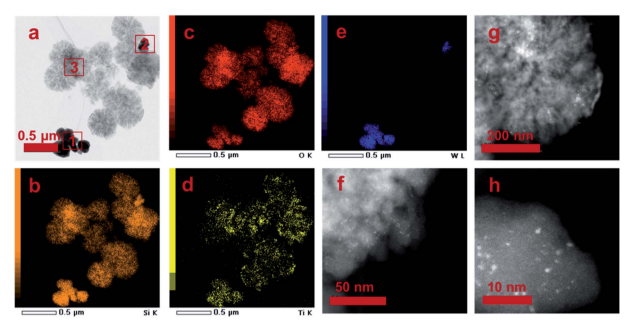


Fig. 5 Microscopic images of WO₃-STS-8: (a) TEM, (b-e) EDS elemental mapping and (f-h) HADDF

Table 3 Element content of different area of WO₃-STS-8

	Content (mol%)					
Element	Fig. 5a (area 1)	Fig. 5a (area 2)	Fig. 5a (area 3)			
0	63.3	58.5	62.9			
Si	4.5	35.6	35.0			
Ti	0.8	3.4	1.6			
W	31.4	2.5	0.5			

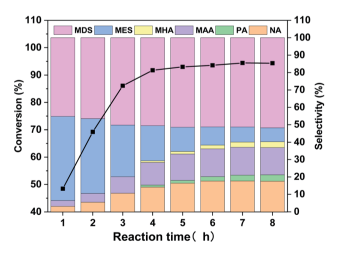


Fig. 6 MO conversion and products distribution along with reaction time for the WO₃-catalyzed MO oxidative cleavage reaction.

selectivity doesn't match the selectivity of NA and MAA in Fig. 6. Because according to Table 4, the selectivity of MDS in Fig. 6 should lower than that of NA or MAA if it is generated from

Table 4 MO Conversion and products selectivity of WO₃-catalyzed oxidation reaction with different starting materials^a

	C	Selectivity (%)					
Starting material	Conversion (%)	NA	PA	MAA	MHA	MES	MDS
МО	91.7	14.2	1.1	13.1	0.9	20.2	50.5
MES	90.0	36.0	0	35.5	0	0	28.5
MDS	0	0	0	0	0	0	0
a n(starting material) : n(H ₂ O ₂) : n(t-BuOH) = 1 : 2.5 : 20, m(WO ₃) : m(starting material) = 0.15 : 10, temperature 80 °C, reaction time 4 h.							

MES. Therefore, most of MDS might be generated directly from MO. As for catalytic reaction of MDS, the result shows that is too stable to occur further reaction under the oxidative reaction condition.

Based on the above results and analysis of WO₃ catalyzed reactions, we speculated the possible reaction pathway of MO oxidation cleavage as depicted in Scheme 1. MO is firstly converted into epoxide (MES) and diol (MDS) products. MDS is relatively inert and cannot go further conversion under the reaction conditions. While MES is reactive and further undergoes hydrolysis to form MDS or oxidative cleavage to form NA and MAA. The aldehydes can be further oxidized to carboxylic acid. As the oxidative cleavage is stoichiometric reaction, the molar ratio of NA to MAA is close to 1:1. As mentioned above, STS is capable of catalyze the oxidation of MO to generate MES, so there would be more MES that could be converted to NA and MAA by WO₃. And that's why the selectivity of NA and MAA is improved and to the opposite, the MDS selectivity is decreased (Table 2, entry 6).

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Scheme 1 Reaction pathway for WO_3 catalyzed MO oxidative cleavage reaction.

The catalytic nature of STS doesn't explain for the improvement of reusability for WO₃-STS in the recycling reaction. The deposition of WO3 on STS might matters. As previously discussed, the catalysis of WO3 on oxidative cleavage of MO is ensured, but how does this happen?. Through the interaction of WO₃ with H₂O₂, peroxotungstate which is the key active species is generated. So MO could be epoxidated and further cleaved into targeted products through the interaction with it. And the peroxotungstate species would turn into polytungstate. Though insoluble in water, it could still react with H₂O₂ and forming the reactive peroxotungstate species and proceeding the MO oxidative cleavage reaction. If H₂O₂ is not depleted completely after the reaction, polytungstate would kept dissolving and generating peroxotung state species which would lead to the loss of tungsten after separation of catalyst from liquid and thus lower down the reusability of the catalyst.

If H₂O₂ is decomposed and consumed, whether it can stop the formation of peroxotungstate species and retain tungsten in solid phase or not?. To verify this, we chose three types of tungsten containing compounds and reacted with H₂O₂ under the reaction condition except MO is absent from it. After the reaction, MnO₂/Al₂O₃(10 w% MnO₂) which was applied for peroxide compounds decomposition in the reaction mixture was added in and we waited until bubbles is not generating obviously. The tungsten concentration in the solution is measured by ICP-OES before and after adding MnO2/Al2O3 and the results are shown in Table 5. It clearly shows that though different in solubility due to the different tungsten containing compound, the concentration of homogeneous tungsten species decreases to lower than 1 mg kg⁻¹. And that is to say, control the content of peroxide compounds in the liquid phase is the key to reduce the loss of tungsten.

The H_2O_2 conversion along with reaction time during the MO oxidative cleavage reaction under different catalyst condition was analyzed by titration method and the results are

depicted in Fig. 7. When no catalyst is employed, the H₂O₂ conversion is lower than 5%. The H₂O₂ conversion increases linearly with reaction time, and reaches to 43% after reaction for 4 h. Compared with WO₃, STS is far more effective for catalyzing H₂O₂ conversion. Since MO conversion is restively low as catalyzed by STS (Table 2, entry 3), part of H₂O₂ in this experiment might be decomposed inefficiently. And this result confirms the peroxide decomposing effect for STS. As WO₃-STS is used, an improved H₂O₂ conversion compared with STS is obtained. As can be seen from the above results, STS functions as epoxidation catalyst as well as peroxide decomposing catalyst.

3.5 Reaction mechanism

The oxidatIve cleavage of MO catalyzed by WO3 and H2O2 is generally considered as homogeneous reaction according to previous studies.31 We prepared homogeneous active tungsten species generated by the reaction of WO_3 and H_2O_2 , and applied it to MO oxidative cleavage reaction with or without STS, as shown in Fig. 8. Firstly, the reaction between WO₃ and H₂O₂ is conducted under 80 °C for 1 h (Fig. 8a). The transparent filtrate is then obtained while it was still hot (Fig. 8b). Part of this filtrate was added with MO and reacted under 80 °C for 6 h. Product analysis shows that MO conversion rate reaches to 91.2%, NA and MAA selectivity is 14.0% and 14.2%, respectively. On the other hand, STS was added to the other part of filtrate reacting for 15 min and the intermediate was obtained (Fig. 8d). Then, MO was added for oxidative cleavage reaction. Product analysis shows that MO conversion rate reaches to 90.5%, NA and MAA selectivity is 28.4% and 29.7%, respectively. As can be seen, the selectivity of NA and MAA has doubled when STS was added in while MO conversion did not decrease obviously. This result again strongly infers that STS was involved in the reaction.

Cold-spray ionization(CSI) mass spectrometer (MS) was further used to analyze the distribution of peroxotungstate species. Four liquid samples of filtrate obtained from the product of Fig. 8b–e were analyzed by CSI-MS. Only two samples from Fig. 8b and d showed peroxotungstate signals as shown in Fig. 9. The signals were attributed to various homogeneous tungsten species, such us $[HWO_4]^-(m/z=249.08)$, $[HWO_3(O_2)]^-$ (m/z=264.88), $[H_{19}WO_{10}(O_2)_3]^-$ (m/z=458.89), $[H_{20}WO_{11}(O_2)_3]^-$ (m/z=476.72), $H_{17}W_2O_{10}(O_2)_5]^-$ (m/z=705.43), $H_{13}W_3O_9(O_2)_6]^-$ (m/z=933.20). When STS was added to the filtrate (Fig. 8d), it was found that the tungsten species with high mass charge ratio disappeared (Fig. 8c). This results indicate that these tungsten species is not stable with the existence of STS, and that explain how the STS promoted the depletion of peroxides.

The reaction mechanism of WO₃-STS catalyzed MO oxidative cleavage is proposed and depicted in Scheme 2. The oxidative cleavage reaction of MO is a two-step process and the epoxide MES is the key intermediate for the whole conversion. To increase the yield of oxidative cleavage products (NA and MAA), it is necessary to enhance the conversion of MO to MES. Both STS and WO₃ are capable of active H₂O₂ and catalyze the expoxidation of MO. In this regard, adding STS into the reaction system would be a good choice due to their excellent catalytic activity for epoxidation.^{34,35} The epoxidation reaction catalyzed

Table 5 Effect of H₂O₂ decomposition on tungsten concentration^a

	Tungsten concentration (mg kg ⁻¹)				
Tungsten containing compounds	Before adding MnO ₂ /Al ₂ O ₃	After adding MnO ₂ /Al ₂ O ₃			
WO_3	138	<1			
$H_{28}N_6O_{41}W_{12}$	384	<1			
$(NH_4)_{10}W_{12}O_{41}\sim xH_2O$	172	<1			
^a $n(\text{Tungsten containing compound}): n(H2O2): n(t_1)$	-BuOH) = 1 : 2.5 : 20, temperature 80 °C.				

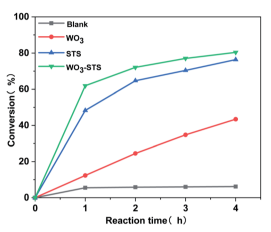


Fig. 7 H_2O_2 conversion along with reaction time catalyzed by different catalyst. $n(MO): n(H_2O_2): n(t-BuOH) = 1: 2.5: 20, m(WO_3-STS): <math>m(MO) = 1: 10$, temperature 80 °C.

by both STS and WO $_3$ supplies sufficient MES for the oxidative reaction. And the selectivity of unwanted diol product MDS which mainly formed directly from MO decreased obviously (Table 2, entry 4 and 6). Furthermore, WO $_3$ is effective in converting MES into NA and MAA and it is the main catalyst to catalyze the MES oxidative cleavage reaction. Thus the binary WO $_3$ –STS catalyst shows high catalytic performance.

Due to the excessive H_2O_2 , insoluble WO_3 kept dissolving and forming soluble peroxotungstate and thus leading to the tungsten leaching. STS is capable of decomposing peroxide, therefore, the WO_3 dissolving process is limited and insoluble tungsten species start to precipitate. When H_2O_2 was completely consumed, the tungstate decomposes into WO_3 particles, which has poorer solubility, and these particles started to aggregate and precipitate. Because of the relatively large specific surface area of STS, it easy to efficiently capture these sub-nano WO_3 particles and recover it. These WO_3 particles are soluble to H_2O_2 and owing to the small size of it, it is easier to be

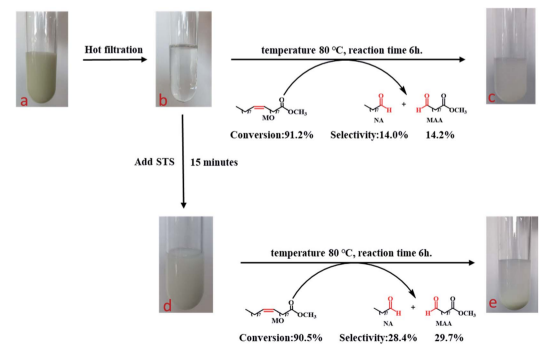


Fig. 8 Hot peroxotungstate-containing filtrate catalyzed MO oxidative cleavage

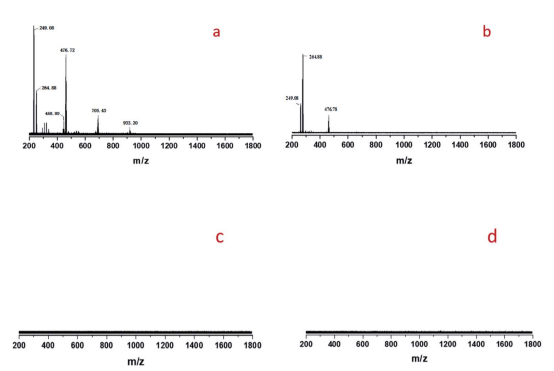
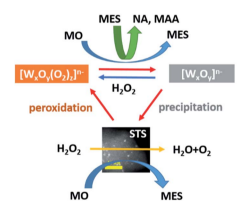


Fig. 9 CSI-MS (negative-ion mode) analysis results of the filtrate: (a) sample obtained from Fig. 8b, (b) sample obtained from Fig. 8d, (c) sample obtained from Fig. 8c, (d) sample obtained from Fig. 8e.



Scheme 2 Proposed reaction mechanism of WO_3 -STS catalyzed MO oxidative cleavage reaction.

peroxidized than bulk WO₃ particles and the MO conversion is thus improved in the recycling reactions. Owing to the cooperation of STS and WO₃, both good catalytic performance and reusability are achieved.

4. Conclusion

In this study, we investigated MO oxidative cleavage reaction catalyzed by WO_3 –STS binary catalyst. Both WO_3 and STS component are capable of catalyze MO epoxidation and generating MES intermediate, while WO_3 is much more efficient in converting MES into targeted products NA and MAA. However, WO_3 is likely to dissolve in H_2O_2 and form soluble peroxotungstate species which results tungsten leaching. STS zeolite

is just able to decompose peroxide and promote precipitation of peroxotungstate species on the surface of STS, so almost all tungsten species can be recovered. Owing to the cooperate catalysis of WO₃ and STS zeolite, the WO₃–STS catalyst shows enhanced catalytic performance with 94.4% MO conversion and oxidative cleavage products selectivity above 63%, and much improved reusability compared with single component of this binary system. These results shed light on designing of highly efficient catalysts for oxidative cleavage of unsaturated fatty acid esters and exhibit great potential for green process developing.

Author contribution

Shihao Xin: investigation, data curation, writing – original draft. Xinxin Peng: conceptualization, methodology, formal analysis, writing – review & editing. Yao Zhang: conceptualization. Aiguo Zheng: tem characterization and investigation. Changjiu Xia: formal analysis. Min Lin: validation. Bin Zhu: validation. Zuoxin Huang: formal analysis, supervision. Xingtian Shu: supervision, project administration.

Conflicts of interest

The authors declare of no conflicts of interest.

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