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# GAS PHASE GLYCEROL DEHYDRATION TO ACROLEIN USING SUPPORTED SILICOTUNGSTIC ACID CATALYST

(Penyahhidratan Gliserol Fasa Gas Kepada Akrolein Menggunakan Sokongan Mangkin Asid Silikotungstik)

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#### Abstract

The gas phase dehydration of glycerol to acrolein over a series of supported silicotungstic acid (HSiW) on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> nanoparticle (W<sub>10</sub>-Al, W<sub>20</sub>-Al, W<sub>30</sub>-Al and W<sub>40</sub>-Al) has been investigated. The catalysts were characterized by temperature programmed desorption, nitrogen adsorption—desorption, thermogravimetric analysis, X-ray diffraction, field-emission scanning electron microscopy and energy dispersive X-ray techniques. The large pore diameters (>21 nm) of the prepared catalysts alleviated the coke deposition effect. The acidity of the samples increased from 2 to 2.6 mmol/g.cat by varying HSiW loadings from 10 to 40 wt.% on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> nanoparticle. The highest acrolein yield achieved was 74.1% at 94% glycerol conversion over W<sub>20</sub>-Al catalyst for 10 wt% glycerol feed concentration and 300 °C reaction temperature in 3 hours. The combined physicochemical characteristics of W<sub>20</sub>-Al made it more superior compared with other samples in the current study.

Keywords: acrolein, dehydration, glycerol, supported silicotungstic acid

#### Abstrak

Penyahhidratan gliserol fasa gas kepada akrolein dengan menggunakan satu siri asid silikotungstik disokong (HSiW) terhadapnanopartikel  $\gamma$ -Al $_2$ O $_3$  (W $_{10}$ -Al, W $_{20}$ -Al, W $_{30}$ -Al, dan W $_{40}$ -Al) telah dikaji. Mangkin-mangkin tersebut dicirikan oleh penyahjerapan suhu terprogram, penjerapan-penyahjerapan nitrogen, analisis termogravimetrik, pembelauan sinar X, mikroskop pengimbasan elektron pancaran medan, dan teknik penyerakan tenaga sinar X. Diameter liang besar (>21 nm) bagi mangkin yang disediakan dapat mengurangkan kesan pemendapan kok. Keasidan sampel meningkat daripada 2 hingga 2.6 mmol/g.cat dengan mengubah beban HSiW daripada 10 hingga 40 wt.% pada nanopartikel  $\gamma$ -Al $_2$ O $_3$ . Hasil akrolein tertinggi yang dicapai adalah 74.1% pada 94% penukaran gliserol terhadap mangkin W $_{20}$ -Al bagi 10 wt.% kepekatan suapan gliserol dan 300 °C suhu tindak balas dalam masa 3 jam. Ciri-ciri gabungan fizikokimia W $_{20}$ -Al menjadikannya lebih unggul berbanding sampel lain dalam kajian semasa.

Kata kunci: akrolein, penyahhidratan, gliserol, asid silikotungstik disokong

#### Introduction

Production of value added chemicals from renewable sources as substitute for conventional petroleum-based processes is a hot issue recently. The huge glycerol availability due to large scale production of biodiesel worldwide provides spectacular opportunities for researchers to evaluate its application as a bio-renewable feedstock in future refineries [1, 2]. In addition, glycerol characteristics such as bio-degradability, non-toxicity, and its multi functional structure made it as one of the top 12 most important bio-based chemicals in the world [3]. Glycerol can be used in

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production of more than 2000 value-added chemicals [4, 5]. One of the most important products which can be produce from glycerol during a bio-based process is acrolein which is a substitute for petroleum based propylene oxidation process [6]. Acrolein is one of the versatile intermediates in chemical industry due to its wide utilization for acrylic acid, superabsorbent polymer, 1,3-propanediol, and many more polymers or polyesters production.

Wide ranges of studies have been reported for application of solid acid catalysts such as heteropoly acids [7-9], zeolites [10, 11], and metal oxides [12-14] in gas or liquid phases of glycerol dehydration to acrolein during the last decade, but still there is no catalyst which exhibit long-term stability without severs deactivation with prospect for industrialization. Among various reported catalysts, supported heteropoly acid (HPA) is attracted much attention due to its stronger acidity (Bronsted acidic sites) and higher activity compared to other catalysts. It is reported that the Bronsted acidic site is more effective than Lewis acidic site for acrolein production [15]. The low specific surface area  $(1-10 \text{ m}^2/\text{g})$  and thermal stability are the main limitations of HPAs. Therefore, to overcome these drawbacks HPAs are often support on a carrier such as alumina, zirconia, and silica. Alumina showed better results due to strong electrical interaction with HPAs and also enhancement of their thermal stability [16]. However, application of  $Al_2O_3$  nanoparticles as a support for HPAs is never reported before. Suitable pore structure and high surface acidity are the key components for higher activity of catalyst for acrolein production [17, 18].

One of the main obstacles for industrialization of bio-based glycerol dehydration to acrolein is fast deactivation of catalyst by coke deposition. For example, the Cs-HPW catalyst is reported as the highest active catalyst in gas phase dehydration of glycerol to acrolein with 98% acrolein selectivity at 100% glycerol conversion, but this catalyst only could last for a few hours and its activity dropped dramatically beyond this time [19]. Some catalyst characteristics such as small pore diameter, very strong acidic sites, and high reaction temperature promote coke deposition [16]. Various researchers proposed different methods for slowing down or eliminating the coke deposition on the catalyst surface such as co-feeding of oxygen or hydrogen. However, this process may increase the probability of explosive conditions or oxidation of products which finally decrease the acrolein selectivity. Besides, some other researchers studied the catalyst structure by addition of noble metals to improve catalyst activity and stability [20, 21]. The main objective of this study is to synthesize a supported HSiW catalyst with  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> nanoparticles for the first time to obtain a catalyst with high surface acidity, large specific surface area, large pore diameter and high thermal stability for increasing glycerol conversion and its selectivity to acrolein.

#### **Materials and Methods**

#### Catalyst preparation

A series of catalysts with various HSiW loadings (10, 20, 30, and 40 wt.%) on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> were prepared by the incipient-wetness impregnation method. In this method, aqueous HSiW solutions were added drop-wise to the  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> nanoparticles support. The slurries were rigorously stirred for 12 h followed by drying at 110 °C for 18 hours. The HSiW-Al<sub>2</sub>O<sub>3</sub> supported catalysts were denoted as W<sub>10</sub>-Al, W<sub>20</sub>-Al, W<sub>30</sub>-Al, and W<sub>40</sub>-Al for 10, 20, 30 and 40 wt.% of HSiW loading, respectively.

#### Characterization

Nitrogen adsorption/desorption at -196 °C was employed to determine the BET specific surface area, pore volume, and average pore diameter (Thermo Scientific, SURFAR). Prior to analysis, the sample was degassed under vacuum at 200 °C for 4 hours for purification and complete removal of adsorbed moisture. The BJH method was used for average pore diameter calculation. Thermogravimetric analysis (TG-DTA) of prepared catalyst was performed using a THERMO TGA instrument in a temperature range of 30 to 800 °C with a ramp rate of 10 K/min under nitrogen flow. The morphology of the samples was observed by Field Emission Scanning Electron Microscopy (FESEM, HITACHI, SU-8000) with the STEM detector. The total acidity of the prepared catalysts was determined by temperature programmed desorption of ammonia (NH<sub>3</sub>-TPD) using the Micrometrics Auto Chem II instrument. A specific amount of catalyst was dried at 200 °C for 1 hour by saturation of NH<sub>3</sub> at 60 °C for 0.5 hour before the physically adsorbed NH<sub>3</sub> was removed by helium purging at 60 °C for 0.5 hour. X-ray diffraction (XRD) was used to determine the structure of the prepared catalysts and to evaluate the degree of crystallinity. The XRD patterns were obtained by D5000 Siemens instrument using Cukα radiation (40 kev, 40 mA). The patterns were scanned in the 2θ range of 10 – 80° (step width 0.05°, 1s per step).

#### Catalytic reaction

The gas phase dehydration of glycerol was conducted under atmospheric pressure in a vertical fixed bed quartz reactor (30 cm length, 11 mm i.d.) using 0.5 – 1 g catalyst sandwich between plugs of glass wool. Prior to reaction, the catalyst was pretreated at 300 °C under nitrogen (N2) flow (1200 ml/h) for 1 hour. Liquid aqueous solution of glycerol (10 wt.%) introduced to the pre-heater (300 °C) by a syringe pump for complete vaporization and then swept into the reactor with 2 ml/h flow rate. Gas hourly speed velocity (GHSV) of the inert carrier gas was 1200 h <sup>1</sup>. The products and unconverted glycerol were condensed in a water-ice-salt bath (-5 °C) and collected each hour for analysis. n-Butanol was added to the condensed products as internal standard. The final solution was analyzed by a gas chromatograph (GC), which equipped with capillary column (DB Wax; 30 m × 0.53 mm × 0.25 µm) and FID detector. To achieve effective product separation, the column was held at 40 °C for 4 minutes before the temperature was ramped up to 200 °C with a rate of 12 K/min. The glycerol conversion, product selectivities and yields are defined by equations (1) to (3):

$$Con_{Gl}(\%) = \frac{M_{Gl, in feed} - M_{Gl, in outlet}}{M_{Gl, in feed}} \times 100\%$$

$$Sel_{N}(\%) = \frac{M_{C in product}}{M_{C in Gl, feed} - M_{C in Gl, outlet}} \times 100\%$$
(2)

$$Sel_N(\%) = \frac{M_{C \text{ in product}}}{M_{C \text{ in Gl food}} - M_{C \text{ in Gl outlet}}} \times 100\%$$
(2)

$$Yield_{N}(\%) = Con_{Gl}(\%) \times Sel_{N}(\%)$$
(3)

where, M<sub>Gl</sub>, M<sub>C</sub>, Con<sub>Gl</sub>, Sel<sub>N</sub>, and Yield<sub>N</sub> are moles of glycerol, moles of carbon for each product, glycerol conversion, selectivity and yield of each product (N), respectively.

#### **Results and Discussion**

#### Surface area and pore structure analysis

The textural characteristics, specific surface area (S<sub>BET</sub>), pore volume (V<sub>P</sub>), average pore diameter (D<sub>P</sub>) of supported HSiW catalysts derived from nitrogen physisorption isotherms are presented in Table 1. The specific surface areas of bulk γ-Al<sub>2</sub>O<sub>3</sub> and HSiW are 143.2 and 8.2 m<sup>2</sup>/g, respectively. By increasing the HSiW loading from 10 to 40 wt.% on the γ-Al<sub>2</sub>O<sub>3</sub>, the surface areas of the prepared catalysts decreased from 140.5 to 76.6 m<sup>2</sup>/g. The reduction of surface area during the catalyst preparation steps was due to the plugging of pores by agglomeration [22], crystallization of active compounds (HSiW), and formation of monolayer HSiW surface with less surface area [23]. Impregnation of HSiW on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> also led to reduction of pore volume from 0.5 to 0.2 cm<sup>3</sup>/g for pure  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> to W<sub>40</sub>-Al. In fact, the highest amount of HSiW loading on alumina diminished approximately 50% of the pore volume. The average pore diameter seems to be not effected and it remained stable with about 21 nm.

Table 1.	Bulk and	l supportec	i samples	s surface area	$(S_{BET})$	), pore vo	lume (	Pv)	), and	pore of	diameter	(Pd)	)
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Catalyst	$S_{BET} (m^2/g)$	V <sub>p</sub> (cm <sup>3</sup> /g)	D <sub>p</sub> (nm)
HSiW	8.2	0.02	10.3
$Al_2O_3$	143.2	0.5	22.5
$W_{10}$ -Al	140.5	0.4	21.0
$W_{20}$ -Al	125.7	0.4	21.5
$W_{30}$ -Al	115.7	0.4	21.1
W <sub>40</sub> -Al	76.6	0.2	21.6

#### Acidity (NH<sub>3</sub>-TPD)

The prepared catalysts were analyzed by temperature programmed desorption of ammonia (NH<sub>3</sub>-TPD) to determine the amount and strength of acidity (Table 2 and Figure 1). The strength of acid sites was divided into three categories of weak (150 – ≤300 °C), medium (300 – ≤450 °C), and strong (>450 °C). The position and area of the TPD peaks are closely related to the acid amount and strength [24]. The total acidity of bulk HSiW and γ-Al<sub>2</sub>O<sub>3</sub> were 1.62 and 1.2 mmol/g.cat, respectively. The HSiW possessed the highest acid strength due to its weak and strong acidic sites compared to the  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> with only weak acidity.

		•			•	-	
Cat	Weak (100	– 300 °C)	Medium (300	0 – 450 °C)	Strong (>	- Total <sup>c</sup>	
	Peak (°C) <sup>a</sup>	Acidity <sup>b</sup>	Peak (°C)	Acidity	Peak (°C)	Acidity	Total
HSiW	131 & 250	0.02	-	-	530	1.6	1.62
$Al_2O_3$	127 & 287	1.2	-	-	-	-	1.2
$W_{10}$ -Al	132	1	300	1	-	-	2
W <sub>20</sub> -Al	137	0.5	301	1.8	-	-	2.3
$W_{30}$ -Al	135	0.5	311	2	-	-	2.5
$W_{40}$ -Al	136	0.6	317	2	-	-	2.6

Table 2. NH<sub>3</sub>-TPD acidity of bulk and supported silicotungstic acid catalysts

The acidity increased significantly from 2 to 2.6 mmol/g.cat by increasing the HSiW loading from 10 to 40 wt.% on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub>. All the samples displayed two peaks at 135 and 300 °C which related to the weak and medium acidic sites. Also, the strength of the medium acid sites was higher than the weak ones. For instance, the W<sub>30</sub>-Al and W<sub>40</sub>-Al approximately have 4 times (2 mmol/g.cat) stronger medium acidic sites than their weak acidic sites (0.5-0.6 mmol/g cat). Also, NH<sub>3</sub> peak positions of weak and medium acidity showed small shifts from 132 – 136 °C and 300 – 317 °C which confirmed increasing the acidic cites strength (Figure 1). The main reason is the electric interaction between HSiW and support ( $\gamma$ -Al<sub>2</sub>O<sub>3</sub>). In fact, during the impregnation process the hydroxyl groups on the surface of  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> were protonated with water which caused a positive charged surface. Finally, the HSiW addition leads an electric interaction. The strength of this interaction depends on the type of support which is stronger for alumina [7, 16].

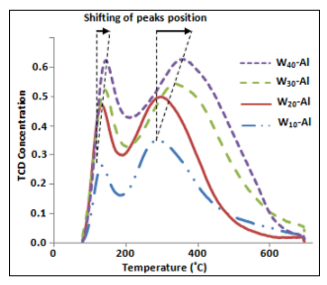


Figure 1. NH<sub>3</sub>-TPD profiles of supported (W<sub>10-40</sub>-Al) samples

#### X-ray diffraction

The X-ray diffraction (XRD) patterns of the supported HSiW (10 - 40 wt.%) on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> nanoparticles are shown in Figure 2. The pure HSiW illustrates three main peaks ( $2\theta = 10.9^{\circ}$ ,  $25.5^{\circ}$ , and  $34.7^{\circ}$ ) which attributed to its Keggin

<sup>&</sup>lt;sup>a</sup> Peak position, <sup>b</sup>Acid amount (NH<sub>3</sub>/Cat, mmol/g), <sup>c</sup> Total acidity (mmol/g.cat)

anions. Besides, the diffraction peaks related to the pure  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> appeared at  $2\theta = 18.75^{\circ}$ ,  $36.9^{\circ}$ ,  $44.2^{\circ}$  and  $67.3^{\circ}$ . However, increasing the HSiW loading from 10 to 40 wt.% on the support caused a reduction in diffraction peaks of  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> and also the reduction magnitude surged. It is noted that none of the supported catalysts illustrates peaks attributed to the HSiW Keggin structure, which demonstrated high dispersion of HSiW on the support. Similar results reported previously [25 – 27]. The main reason for highly uniform dispersion of HSiW on the  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> surface is the size of the Keggin anions (D = 1.2 nm) and the large specific surface area of  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> nanoparticles (S<sub>BET</sub> = 143.2 m<sup>2</sup>/g). Atia et al. [16] mentioned that only >80 wt.% HSiW loading can saturate the support surface. In fact, XRD can detect HSiW crystals whenever high concentration of HSiW saturate the  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> surface and then some HSiW molecules cannot reach the support meso-pores and crystallise as separate phase.

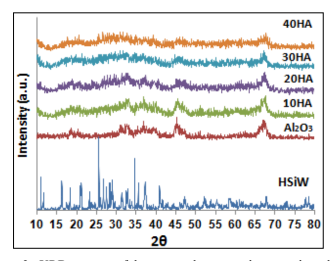


Figure 2. XRD patterns of the pure and supported prepared catalysts

#### Thermal stability

Figure 3 displays the TG-DTA curves of pure  $\gamma$ -Al<sub>2</sub>O<sub>3</sub>, HSiW, and supported W<sub>20</sub>-Al samples. It can clearly be seen that pure  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> lost weight with heating from 30 to 800 °C. The alumina DTA curves exhibits a broad band peak at 70 °C which related to the loss of physically adsorbed water. Afterwards, only a little weight loss observed at 250 °C which attributed to the dehydroxylation of hydroxyl groups. Both TG and DTA curves of the bulk HSiW registered three steps of weight losses. The first two weigh losses of 8% and 3% at 85 and 200 °C, respectively are referred to the loss of physically absorbed water and removal of water from HSiW hydrated heteropoly acid [28, 29]. However, the third weight loss (1%) at 500 °C related to decomposition of the HSiW Keggin anions [16]. Further loading of WO<sub>3</sub> led to new broad band at 3500 cm<sup>-1</sup> crystallite planes, respectively.

Finally the prepared  $W_20$ -Al catalyst shows one broad peak at 80 °C corresponds to the loss of physically adsorbed water (DTA curve). The second weight loss observed at 350 °C which attributed to the removal of structural water molecules. However, no important structural changes were observed in the rest of analysis which confirmed high thermal stability of the  $W_{20}$ -Al sample. In fact, this high stability of  $W_{20}$ -Al catalyst was referred to the electronic interaction between HSiW and  $\gamma$ -Al $_2O_3$  [7, 24]. Scheme 1 depicts different decomposition steps of HSiW and it clearly confirmed that full decomposition happened in >500 °C and the second weight loss of  $W_{20}$ -Alat 350 °C was not the decomposition peak. Remarkably, the prepared catalyst showed high stability even in high temperatures compared to some similar catalysts in previous studies such as HSiW/SBA-15, 30%HSiW-Al, 20%HSiW/A5, 20%HSiW/AS4 which were unstable at temperatures above 480 °C [16, 20, 24].

$$H_{4}SiW_{12}O_{40}nH_{2}O \underset{<200}{\longleftarrow} H_{4}SiW_{12}O_{40} \underset{>500}{\longleftarrow} SiW_{12}O_{38} \rightarrow 12WO_{3} + SiO_{2}$$

Scheme 1. Different steps of silicotungstic acid thermal decomposition

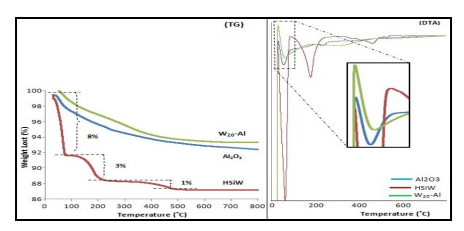


Figure 3. TG-DTA plots for non-supported and supported silicotungstic acid catalysts

#### Morphological analysis

The surface morphology of the fresh and used  $W_{20}$ -Al samples was examined by FESEM (Figure 4). HSiW seems to be dispersed on the external surface of the alumina and filling the voids between the particles with in a gain (Figure 4a-b). As results the specific surface area decreased from 143.2 to 125.7 m²/g for pure  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> and  $W_{20}$ -Al catalyst respectively according to the BET results.Fortunately, no dramatic change in catalyst surface morphology was observed after dehydration reaction (Figure 4c-d). Indeed, FESEM study demonstrated that the catalyst morphological characteristics remain unaffected during the reaction due to high hydrothermal stability of catalyst (confirmed by TGA results), similar results reported elsewhere [30 – 32]. The EDX analysis results related to the fresh  $W_{20}$ -Al sample are exhibited in Figure 4e. The EDX mapping confirms an extremely uniform distribution of alumina (Figure 4f) and tungsten (Figure 4g) elements throughout the  $W_{20}$ -Al catalyst. In addition, the EDX analysis reveals 71.83% and 17.34% weight percentages of Al and HSiW in this catalyst which found to be very close to the theoretical values of 80 and 20 wt% of  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> and HSiW, respectively (Figure 4h).

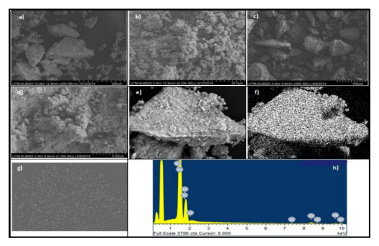


Figure 4. Field emission scanning electron microscopy of fresh and used supported  $W_{20}$ -Al catalyst (a-b), used  $W_{20}$ -Al catalyst (c-d), and EDX mapping analysis of the fresh catalyst: (e) EDX mapping, (f) alumina, (g) tungsten, and (h) EDX mapping result

#### Catalytic performance

Application of bulk samples (HSiW and  $\gamma$ -Al<sub>2</sub>O<sub>3</sub>) in gas phase dehydration of glycerol could not obtain more than 38.1% selectivity of acrolein. These results demonstrate that the strong Lewis acid sites samples ( $\gamma$ -Al<sub>2</sub>O<sub>3</sub>) and even

the strong Bronsted acid site catalyst (HSiW) were not suitable for high acrolein selectivity. Indeed, the strong Lewis acid sites increase the byproducts production and the strong Bronsted acids enhanced coke formation on catalyst surface. Therefore, impregnation of HSiW on support  $(\gamma-Al_2O_3)$  is required to increase the glycerol conversion and acrolein selectivity.

The gas phase dehydration of glycerol to acrolein was investigated by the  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> supported HSiW (10 to 40 wt.%) catalyst at different temperatures (280, 300, and 320 °C) in 3 hours reaction time (Table 3). The results indicated that increasing the HSiW loading from 10 to 20 wt.% led to higher acrolein selectivity but HSiW loading above 20% slightly reduce the selectivity of acrolein or remained constant. Indeed, impregnation of HSiW on  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> caused a significant increase in catalyst activity compared to the pure HSiW due to increase in the total amount of catalytically active protons in the supported catalyst. On the other hand, well dispersed catalyst provides higher accessibility of glycerol molecules to the active compounds of supported catalyst. However, the main reason for reduction of acrolein selectivity or catalyst activity for HSiW loading above 20 wt.% is deposition of a huge amount of active compounds (HSiW) over  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> which cause second surge in the acid strength. Consequently, the coke deposition process on the surface of catalyst increase which has negative effect on acrolein selectivity [33].

The highest acrolein selectivity of 74.1% obtained over the W<sub>20</sub>-Al supported catalyst with 94% glycerol conversion at 300 °C. Also, increasing the temperature from 280 to 320 °C has significant impact on glycerol conversion. The reaction conversion increased from 90% to 97% for 280 to 320 °C, respectively. All the reaction products such as ethanal, propanol, acetic acid, and hydroxyacetone selectivities are summarized in Table 3.

Table 3. Catalytic performance of the bulk and supported (10 to 40 wt.%) silicotungstic acid on nano sized alumina for glycerol dehydration to acrolein<sup>[a]</sup>

Catalyst	T	X	Y	Selectivity (%)							
	(°C)	(%)	(%) <sup>b</sup>	AC <sup>c</sup>	Eth <sup>d</sup>	Pr <sup>e</sup>	Acef	Hyd <sup>g</sup>	Ala <sup>h</sup>	Acai	Total
Blank	300	56	11.6	20.7	-	-	3.8	22.9	-	18.3	65.7
$Al_2O_3$	300	80	30.2	37.7	0.2	0.2	3.8	7.6	2.7	1.2	53.5
HSiW	300	85	32.4	38.1	4.2	5.9	4.5	16.6	5.1	11.2	85.6
	280	89	50.6	56.8	6.9	5.5	4.5	12.5	1.5	3.8	91.5
$W_{10}$ -Al	300	92	55.4	60.2	4.5	4.2	3.4	10.7	1.3	6.1	90.6
	320	93	50.8	54.6	8.7	3.3	3.9	14.1	1.9	6.3	92.7
	280	90	64.2	71.3	2.2	2.1	3.1	3.6	4.1	8.3	94.7
$W_{20}$ -Al	300	94	69.7	74.1	1.1	1.1	2.4	3.3	5.8	5.3	93.2
	320	97	60.7	62.6	3.9	4.9	5.6	5.6	5.5	6.7	94.7
***	280	87	53.2	61.1	4.9	3.6	5.3	7.9	4.1	5.8	92.8
$W_{30}$ -Al	300	91	60.9	66.9	4.1	4.9	5.4	6.1	1.3	6.1	94.8
	320	93	55.6	59.7	5.2	5.9	4.8	6.2	8.5	4.8	95.2
***	280	86	55.8	64.8	5.4	4.3	5.9	5.9	1.5	6.5	94.4
W <sub>40</sub> -Al	300	88	56.3	64.0	5.2	5.2	4.4	7.9	1.8	6.4	94.8
	320	95	58.8	61.9	4.6	4.1	3.6	9.5	1.2	7.2	92.1

<sup>[</sup>a] Reaction condition: Feed: 2 ml/h, 10 wt% aqueous glycerol solution, Carrier gas: 20 ml/min, Reaction temperature: 300°C, Time on Stream (TOS = 3 h), 0.5 g catalyst weight. [b] Acrolein yield, [c] Acrolein, [d] Ethanal, [e] Propanal, [f] Hydroxyacetone, [g] Allyl alcohol, [h] Acetic acid.

The strong interaction between HSiW and  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> is required for long-life stability of catalyst [34]. The TG-DTA results demonstrated the strong binding between HSiW and  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> with high catalyst stability. In addition, the

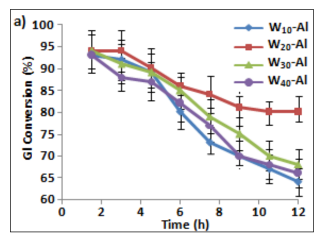
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BET results also indicated that supported  $W_{20}$ -Al catalysts have large pore diameter (> 21nm) which is one of the main reason for high stability of this catalyst. In fact, pore size >4 nm providing a required space for Keggin anions (D = 1.2 nm) and also increase the interaction between adsorbed glycerol molecules and Keggin units [16]. Furthermore, large pore diameter significantly reduces the effect of internal mass transfer limitations [35].

The glycerol conversion related to all the prepared catalysts decreased steadily during 12 hours reaction time. However,  $W_{20}$ -Al recorded the least conversion reduction among all the tested samples (Figure 5a). The acrolein selectivities for all four samples are exhibited in Figure 5b. All samples exhibit dramatic reduction in acrolein selectivity except for the most active and stable catalyst,  $W_{20}$ -Al, which displayed a difference of only 11.6% from 74.1 to 62.5%. The  $W_{20}$ -Al sample was more active than  $W_{10}$ -Al due to the approximately 2 times stronger medium acidic sites (1.8 mmol/g.cat) of  $W_{20}$ -Al compared to the  $W_{10}$ -Al (1 mmol/g.cat). However,  $W_{20}$ -Al only showed 0.2 mmol/g.cat less medium acidity compared to the  $W_{30}$ -Al and  $W_{40}$ -Al samples. In fact, the  $W_{20}$ -Al,  $W_{30}$ -Al and  $W_{40}$ -Al samples showed similar nature of acidity (high medium acid strength). The differences in the total acidity of these samples were only 0.2-0.3 mmol/g.cat.

for  $W_{20}$ -Al was lower than the  $W_{30}$ -Al and  $W_{40}$ -Al samples. The number of acid sites increased from 0.018 to 0.034 mmol/m<sup>2</sup> by surging the HSiW loading from 20 to 40 wt.%. In fact, the  $W_{40}$ -Al sample registered double acid sites amount (0.034 mmol/m<sup>2</sup>) compared to the  $W_{20}$ -Al. Therefore, the main reason for significant reduction of catalyst activity for  $W_{30}$ -Al and  $W_{40}$ -Al samples was higher number of active sites which enhance the coke deposition process on the catalyst surface and deactivate the catalyst faster.

As stated in Table 3 and Figure 5b the acrolein selectivity for  $W_{30}$ -Al and  $W_{40}$ -Al samples were initially high; however, as more coke was deposited on the catalyst surface, the selectivity dwindled during 12 hours of reaction time. Consequently, the optimum amount of HSiW loading was found to be 20 wt.%. For above 20 wt.% HSiW loading, the catalyst activity remained stable in the first hour of reaction, but then its activity and acrolein selectivity declined dramatically (Figure 5a-b). Figure 5c exhibits long-term stability of  $W_{20}$ -Al catalyst during 30 hours reaction time. The stability of  $W_{20}$ -Al sample with 61.8% acrolein selectivity at 78% glycerol conversion prevailed up to 30 hours.



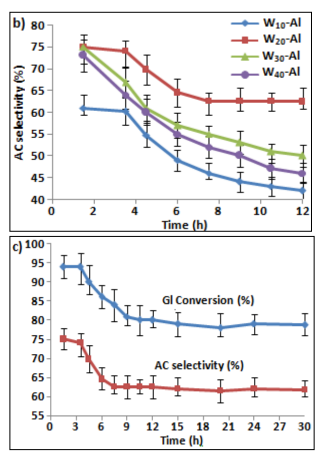


Figure 5. (a) Glycerol conversion versus time and (b) Acrolein selectivity versus time for W10-Al, W20-Al, W30-Al, and W40-Al samples at 300 °C, 12 hours reaction time, 2 ml/h glycerol feed, and 20 ml/min carrier gas flow (c) Long-term stability investigation of W20-Al sample in 30 hours

#### Conclusion

A series of supported silicotungstic acid on aluminum oxide nanoparticle catalysts were prepared and examined in gas phase dehydration of glycerol to acrolein at different temperatures (280, 300, and 320 °C). The  $W_{20}$ -Al catalyst could achieved 74.1% acrolein selectivity with 94% glycerol conversion at optimum reaction condition of 300 °C temperature, 10 wt.% glycerol feed concentration and 0.5 wt.% catalyst amount. The characterization results prove that the  $W_{20}$ -Al large pore diameter (21.5 nm) was the key component for long-life stability of catalyst. In addition, high surface acidity, medium acid cite strength and availability of acid sites on the surface of  $W_{20}$ -Al catalysts led to the highest acrolein selectivity and glycerol conversion among all the synthesized samples.

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