Oxidation of Toluene over LaCrO₃/SiO₂ and LaVO₃/SiO₂ Perovskite catalysts. A comparative study.

H.S.Goswami

Lecturer, Department of Chemistry, Government College, Karauli (Raj.) India

ABSTRACT: Vapour phase catalytic oxidation of toluene over perovskite viz. LaCrO₃/SiO₂ and LaVO₃/SiO₂ has been studied. The heterogeneous catalytic vapour phase oxidation of toluene give benzaldehyde, benzoic acid, maleic acid and CO₂ as products. LaCrO₃/SiO₂ has been found to be the most active and selective catalyst giving 69.3% selectivity of benzaldehyde at 450 °C with surface area 17.1 m²/g. The overall kinetic analysis indicates that the oxidation of toluene to benzaldehyde is first order. The order of catalytic reactivity is LaCrO₃/SiO₂> LaVO₃/SiO₂. The selectivity and activity of catalyst is correlated to surface area value.

KEYWORDS: Heterogeneous Catalysis, Oxidation, Toluene, Benzaldehyde, Surface area.

I. INTRODUCTION

Perovskite type oxides are known to be catalysts for a number of reactions such as total and partial oxidation, hydrocracking, hydrogenation, hydrogenolysis and reduction etc. Amongst the more important reactions in which these compound have been used as catalyst are oxidation of CO^[1-3],CH₄^[4], NH₃^[5], Methanol^[6], Olefins^[7], Paraffin^[8-10], Aromatic compounds^[11-15], Hydrogenation^[16] and oxygenate^[17].

The oxidation of light paraffin's as methane, propane and n-butane has been frequently taken as a test reaction for perovskites oxides. Efforts has largely been directed towards synthesis of unsupported and supported perovskites, oxides of moderates or high specific surface area, their bulk and surface properties and their role in heterogeneous catalysis. The vapour phase oxidation of toluene over mixed oxide has been studied but work with perovskites is scanty. In this note we report the results of a systematic study of LaCrO₃/SiO₂ and LaVO₃/SiO₂ in the oxidation of toluene and compare their reactivity.

II. RELATED WORK

A comparative study on perovskite type mixed oxide catalyst A¹XA₁-XBO₃-λ (A¹=Ca,Sr,A=La,B=Mn,Fe,Co) for NH₃ oxidation was done [5]. Preparation, characterization and catalytic properties of LaMO₃ oxides was done by JMD Tascon et al. Toluene oxidation on LaCoO₃, LaFeO₃ and LaCrO₃ perovskite catalysts was done by K.L.Madhok, D.D. Agarwal and H.S. Goswami.

III. MATERIAL AND METHODS

The LaCrO₃/SiO₂ and LaVO₃/SiO₂ catalyst were prepared by amorphous precursor's citrate decomposition method [18, 19]. Preparation

Supported LaCrO₃ on SiO₂. A general method for preparing 20% LaCrO₃ on Silica is as follows-Aqueous solution of La (NO₃)₃.6H₂O (7.25 gm), Cr (NO₃)₃.6H₂O (6.7 gm), Citric acid (3.5 gm) and Silica (20gm) were mixed in a beaker and then transferred to china dish. The contents were heated at 750°C in a muffle furnace to get the desired catalyst.

Preparing LaVO₃ on SiO₂ are follows: Aqueous solution of La (NO₃)₃.6H₂O (7.27 gm), NH₄VO₃ (1.96 gm), Citric acid (3.45 gm) and Silica (20gm) were mixed in a beaker and then transfer china dish. The final mixture was heated on the sand bath for 10-12Hrs. Finally the catalyst powder so obtained was heated at 600 °C in a muffle furnace to get the desired catalyst.

The surface area of catalyst was determined using ethylene glycol monoethylether (EGME) adsorption method [20,21]. The surface acidity & surface basicity of perovskites catalyst were determined by n-butylamine titration [22] and phenol adsorption method [23] respectively. The I.R. spectra of the perovskites was recorded in KBr using Perkin-Elmer 883 spectrophotometer. The reactor assembly was set up as reported elsewhere [24]. Toluene was taken in a bubbler. The catalyst pellets were heated for 2 hour in a tube furnace at 300 in the stream of air-toluene mixture. Activated catalyst (20 c.c.) was weighed and packed loosely in the reactor. The toluene bubbler preheater zone and furnace zone (catalyst zone) were heated till the required temperature was achieved. The compressed air was allowed to pass through the system.

The air was bubbled through the toluene bubbler and air toluene vapours were passed over the catalyst bed. It was passed through a water condenser. The products were condensed in different containers. Carbon dioxide was estimated from strength of KOH solution. The other products Viz; benzaldehyde, benzoic acid and maleic acid were analysed using literature procedure [25-26].

The oxidation was studied at space velocities (2998 lit hr⁻¹lit⁻¹) temperature (350°C, 400°C, 450°C, 550°C and 600°C) at air toluene ratio (121) respectively.

IV. RESULT AND DISCUSSION

The characterization of the catalyst was carried out using techniques Viz; I.R., Surface area, packing density, surface acidity and surface basicity. The result of these studies has been incorporated in table I and II. The surface area measurements in the temperature range 350°C to 600°C (Table II) shows that surface area of the catalyst increases with increase in temperature up to 450°C but on further increase in temperature the surface area decreases. The specific surface area for different catalyst was found to follow the order LaCrO₃/SiO₂> LaVO₃/SiO₂. The surface acidity and basicity measurements show that LaCrO₃/SiO₂ and LaVO₃/SiO₂ perovskite have acidic and basic sites. The LaCrO₃/SiO₂ is more basic than LaVO₃/SiO₂.

The order of packing density for catalyst is LaVO₃/SiO₂> LaCrO₃/SiO₂.

LaCrO₃/SiO₂ and LaVO₃/SiO₂ were subjected to toluene oxidation. The oxidation of toluene gave benzaldehyde (BzH), benzoic acid (BzA), maleic acid (MA) and CO₂ as the products.

The formation of benzaldehyde as a function of temperature and aerial activity of LaCrO₃/SiO₂ and LaVO₃/SiO₂ at 350°C, 400°C, 450°C, 550°C and 600°C are presented in Table II. The rate of formation of BzH in the beginning is higher on both catalysts. An initial increase in temperature from 350 °C to 450 °C shows an increase in the percentage conversion of BzH from 8.2% to 12.6% on LaCrO₃/SiO₂ (Specific surface area of LaCrO₃/SiO₂ increases from 7.9 m²/g to 17.1 m²/g). On LaVO₃/SiO₂ from 6.4% to 11.2% (Specific surface area of LaVO₃/SiO₂ increases from 10.2 m²/g to 16.4 m²/g).

Further increase in temperature from 450 °C to 600 °C decreases the percent conversion of benzaldehyde (Table II). Thus in the present investigation the catalyst have been found to be active as well as selective at 450 °C for the partial oxidation of toluene. The increase in activity up to 450 °C can be ascribed to increasing removal of trace surface contaminants such as adsorbed gases, hydroxy species of adsorbed water [27] and to the generation of stoichiometric or structural defects such as catalytic site[28]. The decrease in activity observed above 450 °C may be due to a decrease in surface disorder, due to the relatively high mobility of O₂ ions in the lanthanide susquioxides [29] and resulting in the formation of low surface area at 600 °C.

From the result it has been seen that the catalyst is highly select in the oxidation of toluene. The selectivity and activity of catalyst is correlated to surface area values. The most selective and active catalyst is LaCrO₃/Al₂O catalyst because it has more surface area value than LaVO₃/SiO₂ catalyst.

The difference in catalytic activity of these Perovskites heated at different temperature can also be related to the different degrees of heterogenecity of the surface of these oxide [30]. This is caused by terraces, steps, Kinks, Vacancies etc.; having atoms with different degrees of unsaturation and with unusual oxidation states which may play an important role in catalysis.

Based on Haber et at [31]: It can be suggested that toluene activated by the abstraction of hydrogen atom is attacked by a nucleophilic O₂ ion. It can further be suggested that benzaldehyde appears when O₂ ion approaches the – CH₂ group from the direction perpendicular to the benzene ring. This is precisely the direction from which nucleophilic addition of a surface O2 ion of an oxide catalyst could be expected if the toluene molecule were adsorbed side on at an site of the surface through its π electron system. It can thus be concluded that oxidation of toluene to benzaldehyde at the surface of a catalyst is a nucleophilic oxidation.

V. TABLE

Table 1- Characteristics of LaCrO₃/SiO₂ and LaVO₃/SiO₂ Perovskite catalysts.

Catalyst	Decomposition Temperature (°C)	Packing density (g/cc)	Acid st n-butylan me	nine titre,	Base strength n-butylamine titre, meq/q	I.R. frequency
			PKa=6.8 (Neutral red.)	PKa=4.8 (Methyl red.)	Bromo thymol blue PK ₁ =7.2	
LaCrO ₃ /SiO ₂	750 °C	1.24	0.072	0.216	0.65	1080, 800, 780, 695, 600, 460
LaVO ₃ /SiO ₂	600 °C	1.34	0. <mark>016</mark>	0.211	0.63	

Table 2- Activity and Selectivity data.

Catalyst	Surface area m ² /g	Reaction Temp. °C	Conversion % to				Total	%
			BzH	BzA	MA	CO_2	Conversion	selectivity to BzH
LaCrO ₃ /SiO ₂	7.9	350	8.2	2.1	1.9	1.9	14.1	58.1
	10.9	400	9.6	3.2	1.7	1.7	16.2	59.2
	17.1	450	12.6	2.1	1.9	1.7	18.6	69.3
	13.1	550	10.4	3.0	2.0	1.9	17.3	60.1
	12.0	600	9.2	3.1	2.2	2.0	16.5	55.7
	10.2	350	6.4	2.6	2.2	1.2	12.4	51.6
	10.6	400	8.7	2.1	2.9	2.3	16.0	54.3
LaVO ₃ /SiO ₂	16.4	450	11.2	2.1	2.0	1.3	16.6	67.4
	10.4	550	7.4	1.3	3.8	2.6	15.1	49.0
	10.2	600	6.5	4.0	4.0	2.9	17.4	37.3

VI. CONCLUSION

The most selective and active catalyst is LaCrO₃/SiO₂ catalyst at 450 °C for the partial oxidation of toluene. The selectivity and activity of catalyst is correlated to surface area value.

REFERENCES:

- H.R. Khan, H.Frey, J.Alloys compd, 190(2), 209, 1993
- J.M.D. Tascon, L. Gonzalez Tejuca: React, Kinet, Catal.lett, 15, 185, 1980
- H.Faleen, M.J.Martiner-Lope, J.A.Alonso, J.L.G.Fierro solid state Ionics-Elsevier vol. 131, Issues 3-4, page 237-24, 2000 [3]
- G.Landi, P.S.Barbato, A.D.Benedetto, R.Pirone applied catalysis B-environmental Elsevier vol.134-135, page110-122, 2013 [4]
- Yue Wu, Tao yu, Bo sheng, Dou, Cheng-Xian Wang, Xiaofan Xie, Zuo-long yu, Shu-rong Fan, Zhi-Rong Fan, Lian Chi Wang; [5] J.catal, 120, 88, 1989
- [6] B.lavasseur, S.Kalianguine-Applied catalysis A; general Elsevier vol.343 Issues 1-2 page 29-38, 2008
- G.Kremenic, J.M.L.Nieto, J.M.D.Tascon, L.G.Tejuca; J.chem.soc.faraday Trans.1, 81, 939, 1985 [7]
- T.Hayakawa, H.orita, M.Shimizu, K.Takehira, A.G. Anderson, Nomura, Y.Ujihira; Catal.lett, 16(4), 359, 1992 [8]
- [9] H.Arai, T. Yamada, K. Equchi, T. Seiyama; Appl. Catal. 26.265, 1986
- [10] T.Nitadori, S.Kurihara, M.Misono, J.catal, (98) 221, 1986
- [11] Liu, Tuan Chi, Hsu Young Ming; J.chin.inst.chem. Eng. 23(1) 67, 1992
- [12] S.Lars, T.Anderson; J.chem.soc.faraday Trans. 88(1), 83, 1992
- [13] K.L.Madhok; React.kinet.catal.lett., 30,185, 1986.
- [14] S.Irusta, M.P.Pina, M.Menendez, J.Santa Maria journal of catalysis, Elsevier vol. 179, issue 2, page 400-412, 1998
- [15] C.C.Alexandre, Rodrigues; Catalysis Communication Vol. 8, Issue 8, pages 1221-1231, 2007
- [16] S.Amrita Kulkarni, V.Radha Jayaram, Journal of molecular catalysis A, Chemical Elsevier pages 107-110 Volume 223, Issue 1-2, 2004
- [17] T. Shimizu; Appl. Catal; 28,81, 1986
- [18] K.Scott, M.P.Kang, J.Winnick: J Electrochem.Soc; 130, 523, 1983
- [19] R.G.Shetkal 2007 irgu.unigoa.ac.in.
- [20] S.Bhagat, J.D.Ahuja; Ind. J.Chem. 9, 358, 1971
- [21] M.Zawadzki, J.Trawxzynski, Catalysis today Vol. 176, Issue 1, Pages 449-452, 2011
- [22] O.Spalet, J.Balei, I. Paseka; J.Chem.Soe. Faraday Trans. 1,78,2349, 1982
- [23] M.M.Dubinin: In Proc. Int. Symp. On Pre Structure and Properties of materials, Modrys (ed.) Praque, 4, 27, 1973
- [24] K.L.Madhok, K.P.Srivastava, S.Yadav: Indian J. Technol 21,184, 1982
- [25] R. Pereniguez, J.L.Hueso, F.Gaillard, Catalysis Letters, Springer. 142,408-416, 2012
- [26] G.Perchi, M.G.Jilibevto, E.J.Delgado, J.Chem. Technol Biotechnology 86 (8) 1067-73, 2011
- [27] H.Hattori, K.Maruyama, K, Tanabe: J.Catal; 44,50, 1976
- [28] M.P.Rosynek, J.S.Fox: J.Catal; 49,285, 1977
- [29] L.Eyring: In "Handbook of the physics and chemistry of rare earths. Vol. 3, P.337. North Holland, Amsterdam, 1979.
- [30] L.Wachowski, S.Zielinski, A.Burewiez; Acta.Chim.Acad.Sci.Hung; 106,217, 1981
- [31] J.Haber, M.Witko, E.Broclawik: J.Mol. Catal; 45,183, 1980