STUDIES ON THE FORMATION OF MANGANESE MOLYBDATE

P. RAJARAM, B. VISWANATHAN, G. ARAVAMUDAN, V. SRINIVASAN AND M. V. C. SASTRI Department of Chemistry, Indian Institute of Technology, Madras 600036 (India) (Received 2 April 1973)

ABSTRACT

The reaction of MoO₃ with various oxides of manganese (MnO, Mn_2O_3 , Mn_3O_4 and MnO_2) and with $MnCO_3$ has been studied in air and nitrogen atmospheres employing DTA, TG and X-ray diffraction methods, with a view to elucidating the conditions for the formation of $MnMoO_4$. Thermal decomposition of $MnCO_3$ has also been studied in air and nitrogen atmospheres to help understand the mechanism of the reaction between $MnCO_3$ and MoO_3 . The studies reveal that, whereas MnO, Mn_2O_3 and MnO_2 react smoothly with MoO_3 to form $MnMoO_4$, Mn_3O_4 does not react with MoO_3 in the temperature range investigated (480–600 °C). An equimolar mixture of $MnCO_3$ and MoO_3 remains as final product when the same reaction is carried out in nitrogen. Marker studies reveal that manganese ions are the main diffusing species in the reaction between MoO_3 and manganese oxides that result in $MnMoO_4$.

INTRODUCTION

Molybdates of transition metals (AMoO_{α} where A denotes a divalent transition metal ion) offer an interesting class of materials with considerable potential applications. Studies of their structures¹, catalytic properties², infrared absorption³ and magnetic susceptibility⁴ have been reported in recent years. They are usually obtained by reacting MoO₃ with transition metal oxides AO, A₂O₃ or AO₂ at suitable temperatures^{5,6}. Doyle et al.⁷ have reported the formation of MnMoO₄ by heating mixtures of MoO₃ and MnO₂ or MoO₃ and Mn₂O₃ in the required mole proportion in air at 700°C, while Mamykin and Batrakov⁸ have obtained it by heating an equimolar mixture of MnCO₃ and MoO₃ in air in the temperature range 600–930 °C. That only MnMoO₄ (with Mn in +2 state) is obtained as the final product, irrespective of the oxidation state of manganese in the starting material, indicates that the reaction involves an oxidation-reduction mechanism. The present study concerns the mode of formation of MnMoO₄ by reacting MoO₃ with various oxides of manganese (MnO, Mn₂O₃, Mn₃O₄ and MnO₂) and with MnCO₃. Decomposition of MnCO₃ in air and in nitrogen has also been studied with a view to understanding the nature of MnCO₃-MoO₃ reaction.

EXPERIMENTAL

MoO₃ was prepared from ammonium paramolybdate (BDH, AnalaR) by the method given by Palmer⁹. The MnCO₃, MnO₂ and α -Mn₂O₃ used were reagents of certified purity. MnO was obtained by reducing α -Mn₂O₃ in a stream of hydrogen (60 ml/min) at 600°C for 12 h¹⁰.

The differential thermal analysis (DTA) curves of the various solid samples were obtained with Netzsch apparatus (Netzsch Gerätebau, GmbH, Seib, West Germany) while the thermogravimetric (TG) studies were carried out with a Stanton Recording Thermobalance, Model HTSM (Stanton Redcroft Ltd., London). The heating rates for the DTA and TG studies were 10 and 6°C min⁻¹, respectively. For studies in inert atmosphere, cylinder nitrogen was used after passing the gas successively through activated charcoal, magnesium perchlorate, and hot copper gauze $(350-400\degreeC)$.

The solid products obtained at various stages of reaction were analysed by conventional analytical methods¹¹. The total manganese content was determined by EDTA-titration at pH 10 using Eriochrome Black-T indicator, after reduction of manganese in the higher oxidation states to +2 state using hydroxylamine hydrochloride. Higher oxidation states of man_sanese (+3 and +4 states) were determined by sodium oxalate-potassium permanganate method. Molybdenum was determined gravimetrically by the oxinate method.

RESULTS AND DISCUSSION

The reaction, $MnCO_3 + MoO_3 \rightarrow MnMoO_4 + CO_2$, appears deceptively simple. Samples of $MnCO_3$ and MoO_3 heated in air at 400-600 °C for 1-4 h have shown the presence of higher oxidation states of manganese, although prolonged heating gives only $MnMoO_4$. This observation suggests the possibility of an oxidation-reduction mechanism for the formation of $MnMoO_4$. It was thought that a study of the decomposition of $MnCO_3$ itself would provide a clue to the mechanism of $MnCO_3 +$ MoO_3 reaction. The thermal behaviour of $MnCO_3$ was therefore studied in air and inert atmospheres by the methods of DTA and TG and the solid products characterised by chemical analysis and X-ray diffraction. The results (Fig. 1 and Table 1) reveal that $MnCO_3$ decomposes in air at 300-400 °C to yield MnO_2 (endotherm centred around 400 °C in DTA) which subsequently transforms to Mn_2O_3 around 550 °C (endotherm in DTA). Thermogravimetric data show that the Mn_2O_3 formed undergoes further weight loss to yield Mn_3O_4 around 860 °C. These findings on the decomposition of $MnCO_3$ in air are in agreement with those of Paulik and Paulik¹² and Dollimore and Tonge¹³.

The decomposition in inert atmosphere (flowing nitrogen, $\sim 200 \text{ ml/min}$) of MnCO₃ shows only one endothermic peak around 410 °C in the DTA curve. Chemical analysis and X-ray diffraction of the product obtained at this temperature have shown it to be Mn₃O₄. The fact that only Mn₃O₄ is obtained directly from the

decomposition of $MnCO_3$ in inert atmosphere is in conformity with the results of Westerdahl and Leader¹⁴, who too have obtained only Mn_3O_4 , and not MnO as claimed by others^{13,15}, by the decomposition of $MnCO_3$ in argon atmosphere.



Fig. I. DTA and TG curves of MnCO₃ in air and nitrogen.

Atmos- ahere	Temperature range (-C)	Chemical analysis data ?	XRD d spacings of the	Product	° = 11 € 1	ght loss
					Ohs.	Calc.
air	280-400	Mn ^r 63.59% Mn ^{rv} 63.87%	3.11 vs, 2.40 vs, 2.12 s, 1.63 vs, 1.55 s, 1.40 s.	α-MnO ₂	24.30	24.37
	500–580	Mn ^t 69.90% Mn ¹¹¹ 70.04%	3.78 w, 2.71 vs, 2.31 s, 2.00 m, 1.85 s, 1.66 vs, 1.44 w, 1.42 m, 1.38 w.	x-Mn ₂ O ₃	30.70	31.33
	860–890	Mn ^t 71.87%	4.90 w, 3.08 m, 2.87 s, 2.48 vs, 2.37 m, 2.21 s, 2.04 m, 1.79 w, 1.57 m, 1.44 w, 1.34 w, 1.28 w,	Mn₃O⊥	33.20	33.65
nitrogen	320-440	Mn ^t 71.34%	same as above	Mn ₃ O ₄	33.50	33.65

TABLE I

^a Mn^t refers to the total manganese in the solid phase. ^b X-ray diffraction d spacings have been obtained using MoK radiation. The observed d spacings compare very well with those reported for the respective compounds by Moore *et al.*¹⁷. The intensity of the lines is denoted as vs = very strong, s = strong, m = medium and w = weak.

Reaction of MnCO₃ and MoO₃

The DTA and TG results for the reaction between equimolar $MnCO_3$ and MoO_3 mixtures in air are given in Fig. 2 and Table 2. Chemical analyses and X-ray diffraction data for the intermediates and final products are also summarised in Table 2. The DTA results indicate that $MnCO_3$ decomposes around 360–400 °C to yield MnO_2 (endotherm I) which subsequently reacts around 520 °C with MoO_3 forming $MnMoO_4$ (endotherm II). The required weight losses are noted in the corresponding TG curve.



Fig. 2. DTA and TG curves of MnCO₃-MoO₃ mixture in air and nitrogen.

It is to be noted that the formation temperature of MnMoO₄ (~520°C) falls within the range (450–580°C) reported¹³ for the conversion of MnO₂ to α -Mn₂O₃. This strongly suggests, though it cannot be taken as unequivocally proved, that Mn₂O₃ is the reactive intermediate in the reactions of MnCO₃ and of MnO₂ with MoO₃. This is further supported by the finding that mixtures of MnO₂ and MoO₃ and of Mn₂O₃ and MoO₃ in requisite proportions, both lead to the formation of pure MnMoO₄ when heated in air at around 520°C.

The results of the reaction of $MnCO_3$ with MoO_3 in inert atmosphere (flowing N_2) stand in contrast with those obtained in air in that, only a mixture of Mn_3O_4 and MoO_3 , and not $MnMoO_4$, is obtained as the final product when the reaction is carried out in the absence of oxygen. The DTA curve of the mixture (Fig. 2) shows, unlike the curve obtained in air, only one endotherm corresponding to the formation of Mn_3O_4 . That there is no further reaction between Mn_3O_4 and MoO_3 is indicated by the absence of any other heat effects in the DTA curve even up to 700[°]C.

126

Amosphere	System	Temperature	Chemica.	(a21)	XRD d spacings	Product	⁹ /6 IFCIS/I	n loss
		(for Afn a in the pro	nd Mo uduct ^a	tomod sut fo	b3la.tof	Obs,	Calc.
ir	MnCO ₃ -MoO ₃ (1:1)	280-400	Mn ¹ 25 Mn ^{1V} 20 Mo 44	5.12 5.03 77		MnO2 + MoO,	11.20	10.82
		480-600	Muri Muri Muri Muri Muri Muri Muri Muri	1,95 1,95	3.90 m, 3.58 m, 3.44 vs, 3.33 s, 3.21 m, 2.86 m, 2.71 m, 2.51 w, 2.37 w, 2.30 w, 1.99 m, 1.95 m,	MnMaO4	16,70	17,00
	MnO _x -MoO _x (1:1)	480600	Mn ^t 25 Mn ^{tv} 1 Mo 44	5,43 .24 1,28	same as above	MnMoO4	7.12	6.9.3
	Mn2O3-MaO3 (1:2)	500600	Mn ^t 25 Mn ^{tv} 1 Mo 44	5,12 .15 1,61	same as above	MnMo(),	3,52	3,59
itrogen	Mn2O3-M0O3 (1:2)	500600	Mn' 25 Mn ^{Iv} 0 Mo 44	5.34 0.86 1.84	same as above	MnM004	የት	3,59
	MnO _z -MoO _s (1:1)	480-600	Mn ¹ 25 Mn ¹⁹ 0 Mo 44	1,72 1,98 1,72	same as above	MnMoO.	1 07	69,9
	MnCO _a -MoO _a (1:1)	300-440	Mn ^t 25 Mo 44	101 113		$Mn_3O_4 \pm MnO_3$	15,12	14,94

TABLE 2

127

The absence of any reaction between Mn_3O_4 and MoO_3 is further substantiated by heating separate mixtures of Mn_3O_4 and MoO_3 in air and flowing nitrogen at about 600 °C for 2-4 h. In the latter experiments, no formation of $MnMoO_4$ is detected, in contrast to the facile conversion of $MnO_2 + MoO_3$ and $Mn_2O_3 + MoO_3$ to $MnMoO_4$ under similar conditions. Thus, of the various manganese oxides, only Mn_3O_4 appears to be non-reactive towards MoO_3 at least up to 600 °C.

TABLE 3 COMPARATIVE RATES OF FORMATION OF $MnMoG_{\pm}$ FROM $MnCO_3 \pm MoO_3$ AND $MnO_2 \pm MoO_3$ MIXTURES

Temperature	• Time (sec)	Fraction of reaction completed		
		$MnCO_3 + MoO_3$	$\frac{MnO_2 \div MoO_3}{MnO_2 \div MoO_3}$	
500	300	0.848	0.33	
	-400	0.97	0.73	
520	300	0.96	0.74	
	-400	0.98	0.79	
540	300	0.97	0.795	

Isothermal experiments (Table 3)carried out by heating mixtures of oxides of manganese and MoO_3 and of $MnCO_3$ and MoO_3 in the temperature range 500–540 °C, show that $MnMoO_4$ is formed at a faster rate when the carbonate is used as the reactant than when manganese oxides are employed. Apparently the microscopic grain size and poor crystallinity of the freshly formed MnO_2 and Mn_2O_3 are conducive to the diffusion of Mn ions and hence the facile formation of $MnMoO_4$ from $MnCO_3$ and MoO_3 mixtures.

In order to identify the diffusion process that controls the formation of $MnMoO_4$, marker studies¹⁶ were carried out. Pellets of MnO_2 and MoO_3 were pressed together with a thin gold wire marker in between and the composite pellet was heated at 650 C for 30 h. The gold marker was found at the interface of Mn_2O_3 (formed from MnO_2) and $MnMoO_4$. The pellet gave way at the Mn_2O_3 -MnMoO₄ interface while the MoO_3 -MnMoO₄ interface remained intact. The results suggest that manganese ions are the main diffusing species during the formation of MnMoO₄.

ACKNOWLEDGEMENT

This work forms part of a research project sponsored by the National Bureau of Standards, Washington D.C. under contract No. NBS(G)-133.

REFERENCES

1 A. W. Sleight and B. L. Chamberland, Inorg. Chem., 7 (1968) 1672.

2 F. Trifiro and I. Pasquon, Chim. Ind. (Milan), 53 (1971) 577.

- 3 R. G. Brown, J. Denning, A. Hallett and S. D. Ross, Spectrochim. Acta, 26A (1970) 963.
- 4 L. G. Van Uitert, R. C. Sherwood, H. S. Williams, J. J. Robin and W. A. Bonner, J. Phys. Chem. Solids, 25 (1964) 1447.
- 5 R. C. Carlsion, Norelco. Rept., 10 (1963) 8.
- 6 A. P. Young and C. M. Schwartz, Science, 141 (1963) 348.
- 7 W. P. Doyle, G. McGuire and G. M. Clark, J. Inorg. Nucl. Chem., 28 (1966) 1185.
- 8 P. S. Mamykin and N. A. Batrakov, Tr. Ural. Politekh. Inst., 156 (1966) 101: Chem. Abstr., 67 (1968) 39657c.
- 9 W. G. Palmer, Experimental Inorganic Chemistry, Cambridge Univ. Press, London, 1959, p. 406.
- 10 D. G. Klissurski, E. F. McCaffrey and R. A. Ross, Can. J. Chem., 49 (1971) 3778.
- 11 A. I. Vogel, Quantitative Inorganic Analysis, Longmans and Green, London, 1964, pp. 297, 434, 508.
- 12 F. Paulik and J. Paulik, Thermochim. Acta, 3 (1971) 17.
- 13 D. Dollimore and K. H. Tonge, in G.-M. Schwab (Ed.) Reactivity of Solids, Elsevier, Amsterdam, 1965, p. 497.
- 14 R. P. Westerdahl and P. J. Leader, Inorg. Nucl. Chem. Lett., 5 (1969) 199.
- 15 A. J. Hegedus and K. Martin, Microchim. Acta, (1966) 833.
- 16 C. Kooy, in G.-M. Schwab (Ed.), Reactivity of Solids, Elsevier, Amsterdam, 1965, p. 21.
- 17 T. E. Moore, M. Ellis and P. W. Selwood, J. Amer. Chem. Soc., 72 (1950) 856.
- 18 F. Corbet and C. Eyraud, Bull. Soc. Chim. Fr., (1961) 571.