

Thermochimica Acta 255 (1995) 155-170

thermochimica acta

Preparation and thermal dehydration of manganese(II) dicarboxylate hydrates

Yukihiko Suzuki

Department of Applied Chemistry, Faculty of Engineering, Yamagata University, Jonan, Yanezawa 992, Japan

Received 25 July 1994; accepted 1 October 1994

Abstract

Manganese(II) dicarboxylate hydrates $Mn[OOC(CH_2)_n COO] \cdot xH_2O$ have been prepared by the addition of $MnCO_3$ powder or concentrated $MnSO_4$ solution to aqueous solutions of the corresponding dicarboxylic acids.

The crystal forms of the precipitated compounds were observed by optical microscopy. The crystals were obtained either as ellipsoidal, short rods or very small uneven particles. The crystals were different from those of the dicarboxylic acids. The dicarboxylates obtained were characterized by X-ray diffraction analysis and IR spectral measurement.

The thermal dehydrations of the Mn(II) dicarboxylate hydrates were investigated by TG-DTA. The temperatures at which dehydration occurred were taken as a measure of the strength of the Mn-OH₂ bond, and these were found to vary with increasing number of CH₂ groups in the dicarboxylic acid.

The kinetic parameters for the dehydration were calculated by employing a computation method. The three-dimensional diffusion model is found to be the best for describing the kinetic results for the main reaction.

Keywords: Coupled technique; Dehydration; DTA; Kinetics; Manganese dicarboxylate hydrate; TG

1. Introduction

An investigation into the characterization of metal dicarboxylates with both organic and inorganic properties is of great interest because these compounds

0040-6031/95/\$09.50 © 1995 – Elsevier Science B.V. All rights reserved SSDI 0040-6031(94)02132-5

consist of carboxyl groups and metals. Although many investigations of the thermal decomposition reactions of metal dicarboxylates have been carried out [1], they have usually been investigated in isolation, e.g. the decomposition of oxalates and malonates [2], some succinates [3] and fumarates [4].

The effects of systematically varying the cations and keeping the same dicarboxylic anion on the thermal behaviour have been extensively investigated using TG and DTA [2]. There is, however, little or no literature on the thermal behaviour of a series of metal dicarboxylates with the same cation. Manganese(II) ions are labile, can occur in different valence states and will readily change from one valency to another in an air atmosphere. Manganese is fairly electro-positive, and Mn(II) ions form many complexes in which the metal ion is usually octaherdrally coordinated.

Most of these complexes contain no ligand field stabilization energy. The present paper reports the preparation, characterization and dehydration reactions of a series of Mn(II) dicarboxylates using optical microscopy, IR absorption spectrometry, X-ray powder diffractometry, TG and DTA.

2. Experimental

2.1. Purification of dicarboxylic acids

All the carboxylic acids starting materials used were purified by recrystallization from aqueous solutions of reagent grade dicarboxylic acids [HOOC(CH₂)_n COOH]. The crystals obtained were filtered, washed with a mixed solution of ethanol and ether (2:1), and dried over P_2O_5 and then silica gel in a desiccator for several days.

The assay of the acid crystals was determined by titration with standard base.

2.2. Preparation of manganese(II) dicarboxylate hydrates

Mn(II) compounds were prepared by two different procedures.

(1) Oxalate, malonate, azelate, and sebacate were precipitated on reacting $MnCO_3$ powder with an aqueous solution of the corresponding dicarboxylic acids, in 10% excess, at 70°C.

(2) Succinate, glutarate, adipate, pimerate and suberate were obtained by the addition of the corresponding dicarboxylic acid solution to a solution of $MnSO_4$, adjusted to pH 7 with dilute NaOH solution.

In both cases, precipitation of the Mn(II) dicarboxylates occurred immediately. The solution was allowed to stand overnight at room temperature, the crystals were filtered, washed with water and a 2:1 mixture of ethanol and ether, dried by pressing gently between filter papers (Toyo Roshi No. 4), and stored over silica gel in a desiccator. The compounds obtained were pale pink crystalline powders.

Elemental analyses for C, H and Mn are given in Table 1. The number of water molecules depends entirely on the nature of the complex structure and also on the drying procedure [5]. The water of crystallization of the compounds was confirmed

Compound	Formula	C/%		H/%		Mn/%	
		Found	Calcd.	Found	Calcd.	Found	Calcd.
Oxalate	$MnC_2O_4 \cdot 2H_2O$	13.45	13.42	2.24	2.25	30.33	30.70
Malonate	$MnC_3H_2O_4 \cdot 2H_2O$	18.72	18.67	3.09	3.13	28.04	28.46
Succinate	$MnC_4H_4O_4 \cdot 4H_2O$	18.52	19.76	4.27	4.98	21.83	22.60
Glutarate	$MnC_5H_6O_4 \cdot 4H_2O$	23.31	23.36	5.47	5.49	21.32	21.37
Adipate	MnC ₆ H ₈ O ₄ · H ₂ O	33.15	33.20	4.52	4.64	25.28	25.31
Pimelate	$MnC_7H_{10}O_4 \cdot H_2O$	35.85	36.38	5.02	5.23	23.48	23.77
Suberate	$MnC_{8}H_{12}O_{4} \cdot H_{2}O$	39.42	39.20	5.42	5.76	22.40	22.41
Azelate	$MnC_0H_{14}O_4 \cdot H_2O$	42.75	41.71	6.16	6.22	21.91	21.20
Sebacate	$MnC_{10}H_{16}O_4 \cdot H_2O$	44.03	43.97	6.58	6.64	20.02	20.11

Table 1 Analytical data for Mn(II) dicarboxylate hydrates

by elemental and TG analyses. The manganese contents in the dicarboxylates were determined by complexometric titrations with standard EDTA solutions using BT indicator.

2.3. Measurements

TG and DTA curves were obtained using a Seiko Denshi TG/DTA 30 differential microbalance at a heating rate of 10° C min⁻¹ in flowing atmospheres of Ar, CO₂ and air at 200 ml min⁻¹.

About 30 mg of powder sample, in a platinum pan (7 mm in diameter, 70 μ l in volume), was used in each experiment. X-ray powder diffraction analysis and IR spectral measurements were carried out as described in a previous paper [6].

2.4. Kinetic treatment

If the kinetic analysis is carried out with a TG curve obtained at a constant heating rate, $\phi = dT/dt$, and taking into account the Arrhenius equation, the expression

$$g(\alpha) = kt \tag{1}$$

becomes

$$g(\alpha) = \int_0^t k \, \mathrm{d}t = \frac{AE_a}{\phi R} P(x) \tag{2}$$

where $x = E_a/RT$ and $g(\alpha)$ is a function depending on the actual kinetic law. It is based on the quick convergence of the P(x) function, and Eq. (2), after taking logarithms and regrouping terms, can be written as

$$\log \frac{g(\alpha)}{T^2} = \log \frac{AE_a}{\phi R} - \frac{E_a}{2.3RT}$$
(3)

All data were transferred to an NEC 9800 VM microcomputer and linear plots were drawn for the general solid state rate forms [7] of $\log g(\alpha)/T^2$ vs. 1/T by the least-squares method. E_a , A and the corresponding correlation coefficients (γ) for the linear plots were calculated. The high values of the coefficients show a good linearity, and the method which has the best linear relationship, i.e. the coefficient closest to unity, was selected.

3. Results and discussion

3.1. Crystal forms

Typical optical micrographs of the crystals prepared are shown in Fig. 1(a)–(d). The small, uneven crystal forms differed from those of the dicarboxylic acids which have been described in a previous paper [6]. The forms varied with the size of the carboxylic acid group: cubic for oxalate (n = 0), rods for malonate (n = 1), ellipsoids for succinate (n = 2) and adipate (n = 4), and rods for suberate (n = 6) and sebacate (n = 8). The crystals were $80-370 \ \mu m$ in size for the dicarboxylates with an odd number of CH₂ groups, glutarate (n = 3), pimelate (n = 5) and azelate (n = 7), were uneven and the size range was below 30 μm .

3.2. X-ray diffraction (XRD) patterns

The XRD patterns of the Mn(II) dicarboxylate hydrates prepared are shown in Fig. 2(a) - (i). These XRD patterns do not show any similarity indicating that they have different crystal structures. The strongest diffraction lines in the dicarboxylates (acids) appear in a narrow 2θ range, $21.2-25.2^{\circ}$ ($21.4-24.1^{\circ}$), except for 18.3° (29.1°) and 18.5° (23.9°) for oxalate and malonate. The values in parentheses are those of the corresponding dicarboxylic acids. The relationship between the strongest XRD lines and the number of CH₂ groups (n) in the acids and dicarboxylates is shown in Fig. 3. The strongest peaks of dicarboxylates with an even number of CH₂ groups appear in a lower range $21.2-23.6^{\circ}$, while those with an odd number of CH₂ groups are in a higher range, $23.0-25.2^{\circ}$. The values of the strongest XRD lines for the even and odd number of CH₂ groups of the acids and dicarboxylates tend to decrease with increasing n (Fig. 3). In addition, the lines of the acids alternate in character, with higher values for odd numbers and lower values for even numbers of CH₂ groups. However, the lines of the dicarboxylates have lower values with odd numbers and higher values with even numbers of CH_2 groups in the carboxylate unit.

3.3. IR spectra

The IR spectra for Mn(II) dicarboxylate hydrates are shown in Fig. 4, and assignments of the absorption bands are given in Table 2 [8].



Fig. 1. Optical micrographs of (a) $MnC_2O_4 \cdot 2H_2O_3$ (b) $MnC_3H_2O_4 \cdot 2H_2O_3$ (c) $MnC_6H_8O_4 \cdot H_2O_3$ and (d) $MnC_8H_{12}O_4 \cdot H_2O_3$.

In the acids and dicarboxylates, the frequencies of most interest with regard to the structures of the dicarboxylates are those of the C–O vibrations. The IR spectra of the dicarboxylic acids show a v_{asym} (OCO) peak in the vicinity of 1700 cm⁻¹ which is indicative of free carboxyl groups [9]. On complex formation, this v_{asym} (OCO) band is shifted to a lower frequency in all the complexes, showing that complexation is carried out through the carbonyl group [10], and that the O–M bonds in these complexes are essentially electrostatic [11]. Thus it was concluded that the carboxylate group of the acids is coordinated to the Mn(II) ion in the complexes.

Although the $v_{asym}(OCO)$ for all the dicarboxylic acids is approximately constant, for the complexes (n = 6-8) it tends to decrease with increase in n.

The $v_{sym}(OCO)$ band in the complexes was observed between about 1365 cm⁻¹ (0-3) and 1410 cm⁻¹ (4-8). The increase in the difference (Δv) between $v_{asym}(OCO)$ and $v_{sym}(OCO)$ has been taken as a measure of the increasing covalency of the M-O bond [12]. Koppikar and Soundararajan [13] observed that the bidentate coordination of the carboxylate group to the metal results in a lowering of both the v(OCO) frequencies due to the drainage of the electron density from the carboxylate group to the metal; however, at the same time a decrease in the O-C-O angle results in a decrease in separation (Δv). The IR spectra of the Mn(II) dicarboxylates suggest that the bonding of the carboxylate group to the



Fig. 2(a).



Fig. 2. X-ray powder diffraction patterns of Mn(II) dicarboxylate hydrates: (a) oxalate; (b) malonate; (c) succinate; (d) glutarate; (e) adipate; (f) pimelate; (g) suberate; (h) azelate; (i) sebacate.

metal is chelating bidentate in these complexes [14]. The relationship between the separation and n for the acids and the dicarboxylates is shown in Fig. 5. The Δv values for the acids increase with increase in n, while the Δv values for the dicarboxylates decrease with increase in n. The v(C=C) absorptions for the dicarboxylates are lower than the corresponding vibration of the free acids showing that no bonding has taken place with the metal ions (Table 2). The complexes of olefins with Ag fluoroborate [15] and the sandwich complexes of cyclopentadiene [16] gave analogous results. An increase in stability of the complex shifts the stretching vibrations of the C=C bond to lower frequencies.

3.4. Dehydration course

(b)

The TG-DTA curves of Mn(II) dicarboxylate hydrates are shown in Fig. 6. The TG and DTA curves are denoted by solid and broken lines, respectively. Table 3 lists the *n* of the dicarboxylates, the number of dehydration stages, the temperature range of successive dehydrations, the temperature for the peak of the DTA curves

id id	Oxalate	Malonic acid	Malonate	Succinic acid	Succinate	Glutaric acid	Glutarate	Adipic acid	Adipate	Pimelic acid	Pimelate	Suberic acid	Suberate	Azelaic acid	Azelate	Sebacic acid	Sebacate	Assignment
	3380b 1654sb 1620vs 1362m 1362m 1010w 1010w 813vs 813vs	3300 – 2500b 1705m 1314s 1178s	3400b 3350b 1685sb 1660w 1356vs 1356vs 1356vs 1355vs 1352vs 1320vs 858w 858w 858w 859vs 860vs 8720vs	3300 – 2560b 1690s 11425s 1180s	3400b 3150b 2920m 1690m 1650m 11455w 11370vs 11370vs 11390vs 11390vs 11360w 1106w 1106w 1106w 878w 833w 833w 872w 722w	3300 – 2560b 1692s 1300vs 1162s	3300b 3170b 2950m 2950m 1684w 1660w 1676vs 1455m 1455m 1455m 1455m 1455m 1455m 1358m 1358m 1128w 1128w 1128w 1128w 1128w 1109m 934w 934w 934w	3300 – 2500b 1690s 1462m 142m 1432m 1432m 197vs 1143w	3375b 3100b 2916s 2916s 1482m 1482m 1482m 1482m 1482m 1482m 1482m 1133s 133s 133s 133s 133s 1135s 1135	3300 2500b 1465m 1410m 1197s 1197s 11140m 1140m	400b 2960vs 2960vs 1620w 1620w 1660m 1406vs 1340vs	3300 - 2500b 2500b 1470m 1422m 1313m 1313m 1313m 1193s 1193s	3440b 3440b 2970vs 2970vs 1472w 1409s 1353vs 1353vs 1198m 1102w 1020w 1020w 804w 804w 804w 775vs 775vs 772w 712m	3300 - 2500b 2500b 1691s 1432m 1316s 1197s 1137m	3375b 3375b 22990vs 2990vs 1540vs 1466m 1400vs 1332m 1400vs 11193w 1104w 11104w 1104w 1104w 11032w 1002w 770m 770m 770w 770w 770w	3300 – 2500b 1699s 1467m 1430m 1318m 1318m 1318m 1190s	3475b 3475b 2950m 1595sb 1652m 1462m 1462m 1400s 1187w 1187w 1187w 1187w 1187w 750s 7750w 7750w 7750w 7720w 7720w 7720w	(H ₂ O) ((H ₂ O) ((H ₂ O) ((H ₂ O) ((OH) ((OH) ((OH)) ((OH)) ((OH)) ((OH)) ((OH)) ((OH)) ((OH)) ((OC
																		0

Table 2 Infrared absorption data for Mn(II) dicarboxylate hydrates and dicarboxylic acids

Key: b, broad; m, medium; s, strong; v, very; w, weak.



Fig. 3. Relation of the strongest peaks on the X-ray diffraction patterns with the number of CH_2 groups in the Mn(II) dicarboxylate hydrates (solid line) and dicarboxylic acids (broken line).

Table 3 Thermal dehydration data of Mn(II) dicarboxylate hydrates in a flowing Ar atmosphere

Compound	п	Stage	TG temp	$\Delta(t_{\rm i}-t_{\rm e})/$	DTA peak	Weight	loss/%	N'
			range/ C	C	temp ₁ e	Obsd.	Calcd.	
$MnC_2O_4 \cdot 2H_2O$	0	1	116-140	24	134	20.09	20.13	2
MnC ₃ H ₂ O ₄ 2H ₂ O	1	1	128 - 210	82	184	18.52	18.67	2
$MnC_4H_4O_4 \cdot 4H_2O$	2	1	88-136	48	111	28.81	29.64	4
$MnC_5H_6O_4$ $4H_2O$	3	1	67 - 120	53	106	27.24	28.03	4
MnC ₆ H ₈ O ₄ · H ₂ O	4	2	151 - 240	89	191	7.85	8.30	1
$MnC_2H_{10}O_4$ H_2O	5	2	81-119	38	108	7.53	7.80	1
$MnC_8H_{12}O_4 \cdot H_2O$	6	2	120 - 192	72	162	7.28	7.35	1
$MnC_{0}H_{14}O_{4}H_{2}O$	7	2	56-148	92	81	6.50	6.95	1
$MnC_{10}H_{16}O_4 \cdot H_2O$	8	2	65-146	81	98	6.43	6.59	1

Key: N', water molecules lost by dehydration; t_i and t_e , initial and end temperatures for weight loss.



Fig. 4. Infrared spectra of Mn(II) dicarboxylate hydrates: (a) oxalate; (b) malonate; (c) glutarate; (d) adipate; (e) subcrate; (f) sebacate.

 (T_D) , the weight loss after complete dehydration and the number of water molecules lost on dehydration (N'). The dehydration processes in Ar, CO₂ and air for the Mn(II) oxalate to glutarate hydrates (n = 0, 1-3) show a single step, and the Mn(II) adipate to sebacate hydrates (n = 4-8) show two steps. The dehydration behaviour of the dicarboxylates was little affected by the atmosphere, i.e. Ar, CO₂ and air. The observed weight losses for these processes agreed favourably with the calculated water of hydration. The smoothness of the TG and DTA curves and the agreement between the calculated and observed values of weight loss for the dicarboxylates (n = 0, 1-8) suggest that dehydration takes place in similar ways. The endothermic peaks on the DTA curves correlate with loss of water from the crystals. The TG-DTA curves of the dicarboxylates (1-8) in flowing Ar, CO₂ and air atmospheres indicate slow dehydrated rapidly with a somewhat sharp endothermic peak at 134°C. It is possible to suggest that the water of crystallization



Fig. 5. Relation between separation Δv and the number of CH₂ groups in Mn(II) dicarboxylate hydrates (\bigcirc) and dicarboxylic acids (\triangle).

molecules are bonded in two different ways: the water lost at lower temperatures is outer sphere, and the water lost at higher temperatures is coordination water which is strongly bonded with an Mn ion. The initial weight loss temperature (T_i), or the temperature from which the TG curve begins to drop, may be taken as a measure of the relative strengths of the Mn^{m+}-OH₂ bonds. The relationship between T_i and n of the dicarboxylates is given in Fig. 7.

These plots have a similar tendency to those for the strongest peaks on the XRD diagrams and the n of the dicarboxylates described above which were found to change periodically with increasing n.

Heating all the salts at 190° C in the various atmospheres causes a slight decrease in the intensities of the IR absorption bands and XRD signals; at this stage no structural alternation is shown by XRD. From dehydration of the Mn(II) dicarboxylate hydrates, the XRD patterns showed that the crystallinity was somewhat decreased and microscopic observation showed that the particle sizes of the hydrates were subdivided.

Activatio	n energy $E_{\rm a}$ (i	n kcal	mol ⁻¹	an (d freq	luency fact	or A (ir	ו s^-1) fd	or initia	ıl and	main	reaction o	f Mn(II) dicarl	oxyla	tte hyc	drates	in variou	s atmos	oheres
Compound		Stage	Ar						C02		1				Air					
			Initial		Main			Model	Initial	ĺ	Main	1		Model	Initial		Main			Model
			E.	¥	$E_{ m a}$	¥	*		Ē.	٣	$E_{_{u}}$	¥	2		$E_{\rm a}$		Ea	4	4	
Oxalate	(n = 0, X = 2)	Ι	10.1	1.07	67.0	1.37×10^{16}	0.9987	[D3]	11.5	1.15	57.9	7.24×10^{15}	0.9990	LEG I	11 7	901	L C 0L	0201 ~ 11	0.000	
Malonate	(n = 1, X = 2)	I	11.2	2.90	32.6	1.87×10^{9}	0.9981	[D3]	11.2	3.24	31.6	8.26×10^{8}	0.9978	Ĩ		3 0 5	100	1.12×10^{9}	790000	
Succinate	(n = 2, X = 4)	I	9.34	2.65	35.4	1.19×10^{11}	0.9990	[D3]	9.90	2.27	40.1	1.79×10^{12}	0.9986		103	2.2	4 8 24	178×10^{14}	0.0000	
Glutarate	(n = 3, X = 4)	I	9.56	2.30	29.3	3.03×10^{9}	0.9978	[D3]	9.65	1.95	32.0	2.89×10^{10}	0.9983	Ĩ	101	- 19 I		1.70×10^{9}	20000	
Adipate	(n = 4, X = 1)	I	12.3	1.82	45.3	1.31×10^{11}	0.9993	[D3]	11.7	2.70	56.4	1.90×10^{13}	0 9987	5	13.1	5		0.74×10^{12}	0.0000	
Pimelate	(n = 5, X = 1)	1	9.77	1.99	29.0	3.51×10^9	0.9994	[D3]	10.8	1.07	29.5	4.32×10^{9}	0.9984	E G	10.2	1 051	184 8	100×100	0 0000	
		H	11.5	1.67	47.8	1.58×10^{12}	0.9988	[D3]	12.3	1.03	62.8	1.22×10^{15}	1666.0		103	143	35.0 5	01 ~ 1010	0.0070	
Suberate	(n = 6, X = 1)	I	13.6	1.42	32.6	6.81×10^{10}	0.9985	[R2]	14.1	3.40	26.8	3.85×10^{8}	0.9988	[R2]	17.1		33.1 7	$.36 \times 10^{\circ}$	0.9985	[2]
		=	ı	I	ı	1			I	;	Ţ	,	I		I	1				[
Azelate	(n = 7, X = 1)	I	I	I	21.7	9.84×10^{7}	1666.0	[D3]	I	I	41.0	3.57×10^{13}	0.9992	[EC]	I		85	77×10^{7}	0 0080	(D2)
		п	10.5	1.67	38.1	6.63×10^{10}	0.9986	[D3]	10.9	2.09	19.6	6.92×10^{6}	0.9983	[D3]	10.5	1 26 3	1 7 68	24×10^{10}	0.0085	3
Sebacate	(n = 8, X = 1)	I	8.31	2.41	32.0	5.53×10^{10}	0.9979	[D3]	10.2	1.05	29.6	1.29×10^{10}	0.9987	[D3]	10.6	1.27	1 2 60	37×10^{10}	0.0086	
		=	9.48	2.83	26.3	2.98×10^{8}	0.9983	[D3]	10.7	1.17	23.3	5.43×10^7	0.9985	[D3]	10.6	1.12 3	34.9 3	$.31 \times 10^{10}$	0.9979	n E E E
Key: D	3, $F(\alpha) = [1 - 1]$	- (1 – 6	$\chi)^{1/3} l^2$:	R2 /	$E(\alpha) =$	1-0-1	v) 1/2,	correla	tion co	- Hore	1	1 Iraal in A	101			ĺ				

colletation coefficient. " I Kcal IS 4.18 J. 3 2 .



Fig. 6. TG-DTA curves for the thermal dehydration of Mn(II) dicarboxylate hydrates in a flowing Ar atmosphere: (a) $MnC_2O_4 \cdot 2H_2O$; (b) $MnC_3H_2O_4 \cdot 2H_2O$; (c) $MnC_4H_4O_4 \cdot 4H_2O$; (d) $MnC_5H_6O_4 \cdot 4H_2O$; (e) $MnC_6H_8O_4 \cdot H_2O$; (f) $MnC_7H_{10}O_4 \cdot H_2O$; (g) $MnC_8H_{12}O_4 \cdot H_2O$; (h) $MnC_9H_{14}O_4 \cdot H_2O$; (i) $MnC_{10}H_{16}O_4 \cdot H_2O$. TG, solid line; DTA, dotted line.

The diffusion mechanism of the dehydration reaction of Mn(II) dicarboxylate hydrates assumes that the effective area of the reaction interface does not change with the degree of reaction (α), and the rate of evolution of water from the hydrates can be controlled by the rate of three-dimensional diffusion (D3 function) of the appropriate constituent to the active surface. Therefore, it is suggested that the water is loosely accommodated within the channels of the lattice from which escape occurs with minor structural reorganization.

The comparison of fit to the various kinetic equations (those commonly used in kinetics analyses for solid state reaction, namely D1-D4, F1-F3, R1-R3, A2-A3)



Number of CH_2 groups

Fig. 7. Relation between T_i , T_D and the number of CH₂ groups of Mn(II) dicarboxylate hydrates in flowing Ar atmosphere: \bigcirc , initial dehydration temperature (T_i) ; \bigcirc , DTA peak temperature (T_D) .

for the initial and main reactions, see Table 4, shows that the best fit of data of the main dehydration for most of the Mn(II) dicarboxlate hydrates is obtained with the three-dimensional diffusion (D3 function) model. The activation energy E_a and frequency factor A were in good agreement for the kinetic analysis which divided the initial and main reaction on TG curves of the decomposition of dicarboxylic acids as defined in Ref. [17]. The two-dimensional phase boundary reaction R2 was also fitted, but only for Mn(II) suberate. However, when mechanisms other than D3 were selected a somewhat poor correlation coefficient was obtained for the R3 function. The kinetic parameters were calculated by employing a computational method. The computer flow diagram for calculation of Ref. [17]. The relation between activation energy E_a for the initial and main dehydration [17] versus the number of CH₂ groups for the salts is shown in Fig. 8. The plots of Figs. 7 and 8 follow zig-zag lines which is presumed to reflect the zig-zag structure of the dicarboxylic acids.



Number of CH₂ groups

Fig. 8. Relation between activation energy E_a and number of CH₂ groups *n* in the dehydration of Mn(II) dicarboxylate hydrates in flowing Ar atmosphere: \bigcirc and \bullet are initial and main reactions.

Acknowledgment

The author thanks Dr K. Muraishi, Yamagata Women's Junior College, for many helpful discussions and suggestions during this work.

References

 Beilstein Handbuch der Organischen Chemie, Vols. 2-3, Springer, Berlin, 1976, Syst. Nr. 171-179, s. 1877-2202.

Gmelin Handbuch der Anorganischen Chemie, Mn, Koordinations-Verbindungen 2, Springer, Berlin, 1980 Syst. Nr. 56, s. 125-141.

- [2] W.E. Brown, D. Dollimore and A.K. Galwey, in C.H. Banford and C.F.H. Tipper (Eds.), Comprehensive Chemical Kinetics, Vol. 22, Reaction in the Solid State, Elsevier, Amsterdam, 1980, p. 208.
- [3] J.R. Allen, B.R. Carson, D.L. Gerrard and S. Hoey, Thermochim. Acta, 158 (1990) 91.
- [4] J.R. Allen, J.G. Bonner, H.J. Bowley, D.L. Gerrard and S. Hoey, Thermochim. Acta, 141 (1989) 227.

- [5] N.J. Carr and A.K. Galwey, Proc. R. Soc. London Ser. A, 404 (1986) 101.
- [6] Y. Suzuki, K. Muraishi and K. Matsuki, Thermochim. Acta, 211 (1992) 171.
- J.H. Sharp, G.W. Brindley and B.N.N. Achar, J. Am. Ceram. Soc., 49 (1966) 379.
 D. Dollimore, T.A. Evans, Y.F. Lee and F.W. Wilburn, Thermochim. Acta, 188 (1991) 77.
 A.R. Salvador and E.G. Calvo, Thermochim. Acta, 203 (1992) 67.
- [8] K. Muraishi and K. Nagase, Thermochim. Acta, 159 (1990) 225.
- [9] N.B. Colthup, J. Opt. Soc. Am., 40 (1950) 397.
- [10] D.H. Bush and J.G. Bailar, Jr., J. Am. Chem. Soc., 75 (1953) 4574.
 S. Kirrschner, J. Am. Chem. Soc., 78 (1956) 2372.
 P.V. Khadikar and N.S. Sapre, Thermochim. Acta, 128 (1988) 55.
- [11] D.K. Koppikar and S. Soundararajan, Curr. Sci., 45 (1976) 3.
- [12] K. Nakamoto, Y. Morimoto and A.F. Martel, J. Am. Chem. Soc., 83 (1961) 4524. P.V. Khadikar, S.M. Ali and B. Heda, J. Therm. Anal., 30 (1985) 305.
- [13] D.K. Koppikar and S. Soundararajan, Monatsh. Chem., 107 (1976) 105.
- [14] G.B. Deacon and R.J. Phillips, Coord. Chem. Rev., 33 (1980) 227.
- [15] H.W. Quinn, J.S. McIntyre and D.J. Peterson, Can. J. Chem., 43 (1965) 2896.
- [16] E.R. Lindguist and E.D. Nelson, Spectrochim. Acta, 10 (1958) 307.
- [17] K. Muraishi, Y. Suzuki and A. Kikuchi, Thermochim. Acta, 239 (1994) 51.