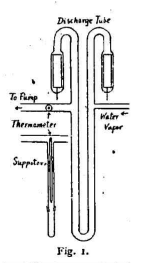
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A COMPARISON OF CATALYTIC ACTIVITIES OF SOME SOLID SALTS IN THE RECOMBINATION OF HYDROGEN FREE ATOMS AND HYDROXYL FREE RADICALS.

By MIKIO TAMURA and SHOH SHIDA.

It is often observed that some salts, such as potassium chloride, have remarkable retarding effects in the combustion reactions of hydrogen and of organic compounds.³⁰ These reactions are chain reactions, and it is usually considered that H-atoms and OH-radicals are chain-carriers in the reactions. Since a retarding action of some negative catalyser is in many cases due to an ability of killing the chain-carriers, it is significant to study the recombination of H-atoms and OH-radicals on the surface of some solid salts.³⁰ In this investigation the relative catalytic activities of some solid salts are determined by measuring the rises in temperature of a thermometer, the mercury-bulb of which was covered with thin layer of solid salt to be tested, and which was inserted into a stream of water vapour involving H and OH. Though the analogous investigation was already made by H. S. Taylor and his collaborators³⁰, the samples used were limited only to several salts. The present author intended to make the same experiment on more samples and more



systematically than they did.

Experimental Method.

The apparatus used is simple and only the main part is shown in Fig. 1. The discharge tube is an ordinary Wood's discharge one. The diameter of the tube is 25mm and the whole length about 1m. The diameter of the reaction tube is also 25mm. The water vapour was generated in a small bulb containing distilled water, and passed through the discharge tube, the reaction tube and a trap cooled with dry-ice and alcohol, and finally pumped out by a mercury diffusion pump and a rotary oil pump. The discharge current was kept constant at 3.0 amp in

1) See for example, N. Semenoff, Chemical kinetics and chain reactions.

2) Three surface reactions involving H-atoms and OH-radicals are possible, that is, $H+H=H_2$, $H+OH=H_2O$ and $OH+OH=H_2O_2$, in which the second is supposed to be most probable.^{*)}

- 3) H. S. Taylor and G. I. Lavin, J. Am. Chem. Soc., 52, 1910 (1930).
- *) Refer to 3) and O. Oldenberg, J. Chem. Phys., 4, 642 (1936).

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the primary of a transformer. A thermometer was inserted perpendicularly into the reaction tube at a place 10cm apart from an exit of the discharge tube. The thermometer, the surface of the mercury-bulb of which was cleaned throughly with NaOH-solution, warm chrom-mixture, water and distilled water, was dipped as far as a constant depth into aqueous solutions of sample salts, the concentrations of which were 1% in most cases. The thermometer was, then, drawn up and dried by heating gently above a soft flame of Bunsen-burner so as to make the deposit to be as homogeneous as possible. After setting the thermometer in the reaction tube, the apparatus was evacuated and then the water vapour flowed through it. After the flowing-rate attained a stationary state, the discharge was started and the temperature rises with times were read. The pressure of water vapour in the tube was 7mm of dibutylphthalate.

Experimental Results.

The experimental results obtained are shown in Tables 1, 2, 3 and 4. Figures in the tables denote the rises in temperature of thermometer.

Exp. No. t (min.)	ĸĘ	z KCl	KBr	4 . KI	<mark>5</mark> ксі	6 NaCl	7 RbCl	8 LiCl
0	25	22	25	25	25	29	26	25
I -	29	48	50 1 94	44	(46	<u> </u>	37
2	32	-	94	(78)4)		-	88	44
3		97	108	· 98			97	_
4	-	104	115	110	96	-	103	-
5	36	107	119	117	103	90	106	
6	<u>u</u>	109	121	I22 .	107.	93	109	· 7
7		117	122	124	110	93	111 -	
8		113	123	126	112	93	112	<u>20</u> 2010
39	4	114	123	127	113	93	115	
10	2 a	114	123	127	115	93	115	(75)
		115		128	115	· ·		
12		С. Х.		128	116			
13					117		10 A	1

Table 1. Series I. Alkali halides 1.

In every series KCl was taken as a standard. The reproducibility of the rise in temperature is satisfactory in the same series, as shown in the experiments on KCl of Series I. and III. For reference the experimental results for Pt were also listed in the table; the reproducibility in the successive runs is almost perfect.

• 4) Gradulations of thermometer between 70~80 could not be read accurately owing to circumstances of the apparatus.

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Exp. No t (min.	, -	I KF		2 KF	R	RbCl			5 KI		6 KBr		7 aC1
o		34		33	-	30			30		30	-	27
I		34		33	8	52	42	8 A	40		54		35
2	1	34		33	s 1		<u>.</u>		(70)		91	- CON	↓ I
3		34		33	0	96	_		94	1	105	1.12	15 .
4		34		33	I	00	93		110		111	0	51
5	ļ	34		33	1	03	95		118		113	-	
6	1	34	t.	33	1	05	97		120		115	8	
7	1	34		32	1	07	98		124		115	6	55
8`		34	2	31	1	07	100		124	2 C	115		
9		34	1	30	1	oS	103		I 24		114	1	
10	ŀ	34	(294)		1	08	106		124		W. 10	6	
лÌ					l		109	2	124		24.0	e l	
12		88.2	100	is i	i		110						
13	•		j.				110		24	1		6	
14					1		111	÷					`
Exp. No. t (min.)	1 NaF	2 LiF	3	Table 4 LigCO3	5	. 6	liscellan 7 NaNO ₃	8	9	IO KNO3	11 MgCl ₂	12 Glass	13 KCl
0	31	33	26	26	26	27	26	30	28	26	32	26	26
` 1	33	31		34	'56	32	56	34	34	6a	35	32	_
2	37	35	65	37		35	So	38	35	117	38	39	
3	39	35	· — ·	38	-	37	102	38	36	131	42	42	100
4	39	35		38	112	37	120	38	36	135	45	42	112
5	38	35	117	37	I 20	37	126	38	36	138	46	42	116
6	'37	35	1 20	36	124.	37	127		365)	141	. 47	40	
7	37	35	s	35	126	35	127			142	-	40	
8	36	35	. 124		128	35	123	•	1	142		39	
9	35	35	125		128	33	.123			141	-		125
10	35		126			33	115			1315)			
		n.,	126		e 8	6	1105)		1	· · · · · · · · · · · · · · · · · · ·	500		

Series V. Carbonates.

Exp. No. . t (min.)	, ксі	Ca(NO3)	CaF	Pt7)	Pt7)	ксі	к,со3	Na ₂ CO ₃
0	• 19	25	26	23	25	23	24	24
. I		29	27	a . <u></u>	162	- 1	} · _`	47
2	92	30	27	240	260	97	1 10	(75)
3	107	29 .	28	306	.320	,100	124	(90)

5) The salts was found after the experiments to have been vapourized.

6) Owing to the hygroscopic property the salt layer did not dry in the atmosphere.

7) Platinized by baking with baking solution.

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	40	12				-		0.25	
	4	113	29	28	350	358	108	136	100
	5	117	29	28	375	377	115	145	112
	6	120	29	28	387	387	115	150	119
	7	121	29	28	391	395	117	153	124
	8	122	28	28	394	395	118	155	127
ŝ	9	122	28	.28	395	396	119	156	129
	10	122	28	23	396	396	120	156	130
3	11	3			396	396	120	1	132
	12				396	396	121		132
		-		1	See	· • • • • • • • • • • • • • • • • •	(he can a	·	

Discusson of Results.

The order of catalytic activeness, which is expressed by the temperature at $t=10^{6}$, is as follows.

 $\begin{aligned} &Pt > (K)^{0} > (Na)^{0} > K_{2}CO_{3} > (KNO_{3})^{0} > KI > KBr > Na_{2}CO_{3} > (NaNO_{3})^{0} > RbCl > KCl \\ &> NaCl > LiCl. \end{aligned}$

LiF, NaF, KF, CaF_2 ; Na₂SO₄, K₂SO₄; NH₄Cl, Ca(NO₃)₂, MgCl₂ have no appreciable catalytic activity.

In these results the regularities of the catalytic activities are found as follows.

- (1) $Rb \rightarrow K \rightarrow Na \rightarrow Li$ (for carbonates, halides and nitrates).
- (2) Metal>Carbonates>(Nitrates)⁹⁾> Halides (for K- and Na-salts)
- (3) -I > -Br > -CI > -F (for K-salts)
- (4) Fluorides and Sulphates are inactive. (for K- and Na-salts)
- (5) Salts of alkali-earth metals are inactive, so far as the present experiments are concerned.

It is concluded that in general the more positive the positive ion is and the less negative the negative ion is, the more active the salt is.

The results showing the higher activities of potassium salts than of sodium salts were expected from the fact that the same tendency as in a negative catalyser had also found in some combustion reactions.

In the experiments on the nitrates, as seen in Series III. No. 7 and 10, after attaining maximum values, the temperature began to fall. This may be due to the reduction of the salt to metal and successive evaporation of the metal by the heat of recombination.¹⁰⁾ Thus, the maximum values obtained for these nitrates are those of the evaporating metals. The values of metals themselves would be

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⁸⁾ Precisely, it should be expressed by the rise in temperature above room-temperature and not above that at t=0, because the latter is not necessarily equal to room-temperature.

⁹⁾ See later.

¹⁰⁾ Indeed, the reduction of nitrates by atomic hydrogen was found by H. Kroepelin and E. Vogel, Z. anorg. Chem., 229, 1 (1936).

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obtained by the extrapolations of the falling curves after the maximums to t=0. The present data, however, are unfortunately not so accurate as to make it possible, and we are obliged to be satisfied with the results that K-metal¹⁰ and Na-metal next to it, are more active than any other salts of them.

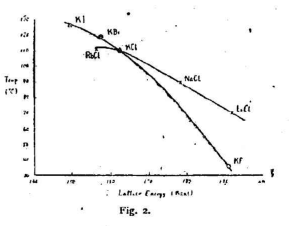
For alkali halides, seeking for the relations between the observed order of activeness and other physical constants, we have arrived at a conclusion that, as shown in Table 5, the smaller the lattice energy is, the greater the catalytic activity is.¹²)

	Salts	Mean value of Temp. at $t=10$	Lattice cnergy ¹⁸ (Kcal)
e e	KF	35	191
Dependence on halogen .	KCl	011	161
ions _	KBr	119	158
• · · · ·	KI	126	158 148
	LICI	70	. 194
Dependence on alkali	NaCl	90	178
ions,	, KCl	110	162
27	RbCl .	111	· 156

the second s	
Table	5.

In an investigation on the influence of alkali halide on, the lowest temperature at which a reaction between methane and oxygen is observed, Jorrissen and Lebbink¹⁴⁰ found that the lowest temperature rose in order Li <Na <K <Rb and similary in order F < Cl < Br < I. These orders agree with those of the catalytic activity in the reactions involving H and

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11) It is noticed that K-metal is well-known as one of the most effective antiknocking substances. 12) If the temperatures are plotted against the lattice energies (Fig. 2), it is seen that, at first sight, the former increases linearly as the latter does with the exception of the point for KF and the slight deviations of the points for KBr and RbCl. However, the deviations of the points for KBr and RbCl cannot be taken as experimental errors. If we take these points as correct ones and take into account the point for KF also, we find that the points for KJ, KBr, KCl and KF fall on one curve and, those for RbCl, KCl, NaCl and LiCl fall on another curve. Therefore, the catalytic activity and the lattice energy are not in a relation of one to one correspondence. In respect to this point more investigations are desired.

13) E.A. Moelwyn-Hughes, Physical Chemistry, P. 601.

14) W.P. Jorissen and H.O. Lebbink, Rec. Trav. Chim., 58, 959 (1939).

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OH in the present investigation. In Table 5 it is seen that the influence of change in halogen ion on the catalytic activity of alkali halide is more pronounced than that of change in alkali ion; similar tendency is also found in the reaction between methane and oxygen. From these facts it is conceivable that the reactions occurring on the surface of alkali halide crystals are the same in both cases. It appears, therefore, that the retarding action of alkali halides in the reaction between methane and oxygen is ascribed to their action of promoting recombination processes involving H and OH.

As seen in the cases of -I, -Br, -Cl and -F, in general, the salt of weaker acid is more active; the high activeness of carbonates and the inactiveness of sulphate may obey this rule.

The authors wish to express their hearty thanks to Professor S. Horiba in the Kyoto Imperial University for his encouragement throughout this investigation.

The Laboratory of Physical Chemistry of the Tokyo Institute of Technology.

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