CARBOXYL GROUP -CO2 EXCHANGE IN THE THERMAL DECOMPOSITION OF ALIPHATIC DICARBOXYLIC ACID SALTS.

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into the residual phenylacetic acid.

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SUMMARY

The exchange reaction of carboxyl groups with carbon dioxide has been studied in thermal decomposition of aliphatic dicarboxylic acid disodium salts. It has been found that exchange takes place under given experimental conditions. In some cases the process results in considerable exchange even in lower conversion ranges therefore it is of significance in the field of isotopes techniques.

1 NERCHUCETON

the mass of aliphatic carboxyl groups in the thermal transformation of aliphatic carboxylate salts can be acted by isotope techniques (Fig.1). The rate of this exchange is known to depend on the nature of corresponding cation. If the exchange process shows some analogy to the dendel reaction then on the basis of a widely accepted mechanism for such transformations (6-3) (Fig.2), exchange between carboxyl groups and $\rm CO_2$ can be expected too. These findings seem to justify the study of this phenomenon in the case of aliphatic carboxylate salts in spite of the observation of Bühler and coworkers (10) who stated that the thermal decomposition process of lithium phenylacetate effected in the presence of $\rm ^{14}CO_2$ was accompanied only with a slight incorporation of $\rm ^{14}C$

Fig.1

In this paper we wish to report our experimental results obtained for the exchange reactions of carboxyl groups with \$1400_2\$ studied in the thermal transformation process of some aliphatic dicarboxylic acid disodium salts.

Fig.2

EXPERIMENTAL

Starting materials

Preparation of sodium salts was carried out by the following method. An equivalent amount of Mach solution was added to curre-oxylic acid in water. The solution was evaporated and curied in vacuo at 200 °C for 2 hours. Purity of the product was checked by Na analysis.

Ethane-1,1,2-tricarboxylic acid

The compound was prepared from ethyl \(\mathcal{J}\)-bromoacetate and diethylmalonate by the method used for pentane-1,1,5-tricarb-oxylic acid. The ester formed was distilled in vacuo (b.f. 155 °C/8 mm Hg). The tricarboxylic acid obtained in the course of alkaline hydrolysis was crystallized from water; m.p. 158-9 °C; yield 52 % (for ethyl \(\mathcal{J}\)-bromoacetate).

Propane-1,1,3-tricarboxylic acid

The triethyl ester (b.p. 155 °C/6 mm ag) was prepared as

acid ethyl ester than subjected to alkaline hydrolisis. The acidified reaction mixture was evaporated to dryness and the tricarboxylic acid dissolved with alcohol from the solid residue. The alcohol was distilled and, on the addition of acetone, the syrupy residue crystallized. Yield 55 % (for ethyl \$\beta\$-bromo-progionate); m.p. 140-42 OC.

Butane-1,1,4-tricarboxylic acid

The triethyl ester of this acid (b.p. 175 °C/12 mm Hg) was prepared and then hydrolyzed by the method mentioned above. The butanetricarboxylic acid was precipitated from concentrated ethyl acetate solution with petroleum ether. Yield 61 % (based on ethyl %-bromobutyrete); m.p. 139-40 °C.

Pentane-1,1,5-tricarboxylic acid

Lthyl **6**-bromovalerate was reacted with diethyl malonate in the presence of sodium ethoxide. The resulting triethyl ester was distilled in vacuo (b.p. 165-70 °C/4 mm Hg) and hydrolyzed in aqueous solution of KOH⁽¹¹⁾. The homogenous solution was concentrated then acidified with cc. HCl to pH l and extracted with ether. The extract was dried over Na₂SO₄ and evaporated to dryness. field 62 % (based on ethyl **6**-bromovalerate); m.p. 39-90 °C (from ether — petroleum ether).

Method for the study of the exchange reaction

The study of thermostability and the choice of reaction conditions were effected on the basis of derivatograms recorded by a n.CM type hardey-Faulik derivatograph in N_2 stream (15 l/hr). The results are shown in Table I.

The exchange reaction carried out in a metal bath thermostat was studied in the following way.

On placing dicarboxylic acid disodium salt (10 mM) in a tube and distilling ¹⁴CO₂ (2 mM; spec.act. 1.88-10⁷ dpm/mM) the tube was sealed and placed in a thermostat and kept at an appropriate temperature. In elaborating the samples, the conversion was determined by the isotope dilution method from an aliquot of the aqueous salt solution. The recovered acids were purified by repeated recrystallizations. Purity of the products was checked

by m.p. and C, H analysis. Radioactivity measurements were performed in dioxane solution by means of a Packard Tri-Carb scintillation spectrometer Model 574. The results are given in Table II.

	Table	I			
Thermogravimetric	parameters	οſ	the	sodium	carboxylates

Salts [¥] _	Decomposition	temperature ⁰ C)	.eight loss (%) in two stage	
	Initial	DTG peak	1	2
MNa	320	350 , 510	12	13
MMNa	370	400,500	18	9
EMNa	370	430, 510	22	10
SNa	420	430 -	33	-
ETNa	330	390 , 470	10	10
GNa	450	500 –	38	-
PRTNa	350	380 , 480	8	14
ANa	440	480 -	41	-
BTNa	400	460 -	32	-
PNa	410	500 -	44	-
PTNa	380	430,500	8	28

Abbreviations:

MNa: Disodium malonate; MaNa: Disodium methylmalonate; EMNa: Disodium ethylmalonate; SNa: Disodium succinate; ETNa: Ethane-1,1,2-tricarboxylic acid trisodium salt;

GNa: Disodium glutarate; PRTNa: Propane-1,1,3-tricarboxylic acid trisodium salt; ANa: Disodium adipate; ETNa: Butane-1,1,4-tricarboxylic acid trisodium salt; Twa: Disodium pimelate; PTNa: Pentane-1,1,5-tricarboxylic acid trisodium salt.

23.4

0

18.8

40.4

н..а

FNa

FNa

PNa

oalts [≭]	Temperature	lime	Conversion	Radioactivity of residue	Exchange**
	(°3)	(min)	(%)	acid (10 ⁶ dpm/mM)	(%)
iña	930	60	53	0.728	21.3
11.Na	<i>5</i> 60	120	72	1.250	36.6
LLVa	580	120	67	0.964	28.2
Sila	440	30	47	2.620	76.6
GNa	420	60	1 5	0.191	5.6
ANa	420	60	0	0	0
ANa	440	60	72	0.711	20.8

Table II Exchange data

60

60

120

120

450

390

410

420

EXFERIMENTAL RESULTS AND DISCUSSION

85

0

40

59

0.300

0.643

1.380

Ω

According to Table II, considerable exchange can be observed between the carboxyl groups and ${\rm CO}_2$ in the thermal decomposition of the salts. This exchange reaction is in some instances so fast that it nearly leads to equilibrium distribution even in lower ranges of conversion, therefore it is of considerable importance in the field of isotope tracer techniques $^{(12)}$.

The remarkable difference in the thermal transformation processes of aromatic and aliphatic carboxylic acid salts may probably be attributed to the presence and great mobility of d-hydrogens $^{(1,j)}$. Consequently, in the course of the thermal accomposition of aliphatic carboxylic salts, hydrogen splitting and subsequent carboxylation will mainly occur on the d-carbon atom resulting malonate type intermediates (Fig. 3).

^{*} Abbreviations are the same as in Table I.

^{**} These particulars give the extent of equilibrium distribution in percentage.

$$R-CH_{2}COOM \longrightarrow (R-CH_{2}^{-}) + CO_{2} + L^{+}$$

$$R-CH_{2}COOM + R-CH_{2}^{-}) \longrightarrow R-CH_{3} + (R-CH-COOM)$$

$$(R-CH-COOM) + CO_{2} + L^{+} \longrightarrow R-CH(COOM)_{2}$$

Fig.3

The latter have, however, shown no thermostability under the experimental conditions chosen (Table I), and after decarboxylation and subsequent hydrogen splitting these are transformed into the starting salts, producing another proton deficient fragment. Thus, in addition to the hydrogen transfer process (14,15), decarboxylation of the malonate type product formed in carboxylation also takes part in this fragment-reproducing process (Fig.4).

$$R-CH$$
 $R-CH^{-14}COONa + CO_2 + Na^+$
 $R^*-CH_2-COONa + R-CH_2^{-14}COONa$
 $R^*-CH_2-COONa + R-CH_2^{-14}COONa$
 $R^*-CH_2-COONa + R-CH_2^{-14}COONa$

It is in the latter reaction that the exchange between carboxyl groups and CO2 occurs in our interpretation of the mechanism.

In our opinion, it is highly probable that the proton deficient fragment in \$\mathcal{J}\$-position plays a decisive role in the exchange reactions. The individual stabilization and formation processes of these fragments constitute the dynamic state of the substance, which, under given experimental condiditions, gives the above fragments a macroscopic existence on the one hand, and, on the other, brings about the above outlined detectable changes.

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