Thermal Decomposition of Aromatic and Heteroaromatic Amino-acids†

By G. P. Shulman* and P. G. Simmonds

(Jet Propulsion Laboratory, California Institute of Technology, 4800 Oak Grove Drive, Pasadena, California 91103)

RECENTLY, investigations of the pyrolysis of amino-acids1,2 or their sodium salts3 have been reported. The method is potentially useful for rapid identification of amino-acid constituents of peptides or proteins. It was apparent that either the analytical techniques (gas chromatography and mass spectroscopy) were inadequate to detect all the products, or that markedly different reaction schemes applied to the closely related amino-acids phenylalanine, tyrosine, tryptophan, and histidine. We have now characterized the volatile products obtained from pyrolysis of these amino-acids, and conclude that there is no significant variation in the mode of decomposition.

A complete gas chromatographic analysis of the phenylalanine pyrolysate is shown in the Figure. The magnitude of quantitative variations (Table) resulting from changes in construction materials are also typical of those obtained by varying pyrolysis temperature from 400 to 600° or pyrolysis time from 4 sec. to 1 min. It is obvious from the products that suitable conditions for oxidation-reduction reactions are established during pyrolysis. Products can be explained as resulting from the following reaction sequences:

$$\begin{array}{c} \operatorname{NH}_{2} & -\operatorname{CO}_{3} & \operatorname{Ph}[\operatorname{CH}_{2}]_{2} \cdot \operatorname{NH}_{2} \\ & & & & & \\ \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} \xrightarrow{-\operatorname{CO}_{3}} & \operatorname{Ph}[\operatorname{CH}_{2}]_{2} \cdot \operatorname{NH}_{2} \\ & & & & & \\ \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} \xrightarrow{-\operatorname{NH}_{3}} & \operatorname{Ph}\operatorname{CH} = \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} \\ & & & & \\ \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} \xrightarrow{-\operatorname{CO}_{2}} & \operatorname{Ph}\operatorname{CH} = \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} \\ & & & & \\ \operatorname{Ph}\operatorname{CH} = \operatorname{CH}_{2} & \operatorname{Ph}\operatorname{CH} = \operatorname{CH}\operatorname{Me} \\ & & & & \\ \operatorname{Ph}\operatorname{CH} = \operatorname{CH}_{2} & \operatorname{Ph}\operatorname{CH} = \operatorname{CH}\operatorname{Me} \\ & & & & \\ \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} & \xrightarrow{-\operatorname{Ph}} & \operatorname{Ph}\operatorname{Ph}\operatorname{Ph}^{n} \\ & & & \\ \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{CH} \cdot \operatorname{CO}_{2}\operatorname{H} & \xrightarrow{-\operatorname{Ph}} & \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{+}\operatorname{CO}_{2} \\ & & & & \\ \operatorname{Ph}\operatorname{CH}_{2} \cdot \operatorname{CO}_{2}\operatorname{H} & \xrightarrow{-\operatorname{Ph}} & \operatorname{Me}\operatorname{CN} \\ \end{array}$$

[†] This paper presents the results of one phase of research carried out at the Jet Propulsion Laboratory, California Institute of Technology, sponsored by the National Aeronautics and Space Administration. Presented at the American Chemical Society, Western Regional Meeting, Anaheim, California, November, 1967.

				Analytical a	lata fro	m þyrol	ysis of an	rino-aci	ids, RC	CH₂·CH	Analytical data from pyrolysis of amino-acids, RCH ₂ ·CH(NH ₂)·CO ₂ H			
Я	0	G.c. column	Pyrolysis temp.	Chamber material	CO_2^{a}	H_2O	CO ₂ ª H ₂ O CH ₃ CN	КH	RMe	Relati REt	Relative percentage of REt RCH=CH ₂ R(of RCH ₂ CN	Relative percentage of REt RCH=CH ₂ RCH ₂ CN RCH ₂ CH ₂ NH ₂ RCH ₂ CH ₂ R	RCH ₂ CH ₂ R
Рћ	:	qΡ	500°	Stainless	Qual	itative	Qualitative analysis							
		B°	400	Stainless	q	р			28	0.74	0.95	8·2	49	6.9
		В	500	Stainless	q	q	1	}	30	6.79	1.1	8·6	53	6.2
		В	600	Stainless	q	q	1		24	0.47	16-0	6.2	64	5.2
		В	500	Stainless	27	20	1.0	2.7	14	0.6	0.6	$5 \cdot 1$	26	1.5
		В	500	Quartz	29	24	0·8	l∙4	28	0·8	1.2	2.5	10	0.5
		В	500	Pyrex	28	19	0·8	1.6	19	0.5	0.4	5.0	21	2.0
		В	500	Pyrex ^e	p	q	ł		44	1.9	4-9	8·2	30	4 ·3
		ü	500	Stainless	Qual	itative	Qualitative analysis							
p-C ₆ H ₄ ·OH	:	V	500	Stainless	Qual	itative	Qualitative analysis							
		В	500	Stainless	52	30	1.0	6	9	63				
Indol-3-yl	:	ပ	500	Stainless	22	12	0-7	27	33					
Imidazolyl	:	D۴	500	Stainless	Qua	litative	Qualitative analysis							
 Including CH₄ and other hyd b Squalane-50 ft. of 0.02 i.d.; Carbowax 20M-200 ft. of 0.4 d Not Adacmined (45 mb ioni) 	lg CH ₄ le—50 lx 20M	and oth ft. of 0-0 [200 ft.	^a Including CH ₄ and other hydrocarbons. ^b Squalane—50 ft. of 0.02 i.d. support coated o ^c Carbowax 20M—200 ft. 06 0.02 i.d. stainless s ^d Not Adverningd (Hound indicator)	ns. coated open : ainless steel :	tubular capillar	columi y coate	a program d with 10	ımed 5(% solu) to 12 tion, p	5° at 6 rogram	° min1 then imed 50 to 20	maintainec 0° at 6° mi	drocarbons. o2 i.d. stainless steel capillary coated with 10% solution, programmed 50 to 200° at 6° min1, then maintained at 200°. 	ained at 200°.
177 3017 H	MITTIN TOA	MITOTI DO	IN TION BOTTON D	creeres /										

 Not determined (name ionization detector).
 Sealed tube—1 min. pyrolysis.
 Sealed tube—2 not 0.2 i.d. stainless steel capillary coated with DC200-Igcpal 880/20:1, programmed 50 to 200° at 4° min., then maintained at 200°. & Ethylene glycol adipate-6 ft. of 4 mm. i.d. glass tubing containing 10% EGA on 80-100 mesh Gas-Chrom Q support.

CHEMICAL COMMUNICATIONS, 1968

An amide formed by condensation of two aminoacids is a likely, though unproven, intermediate products o- or m-cresol, xylenol, and o- or methylphenol. Similarly, tryptophan gave indole,

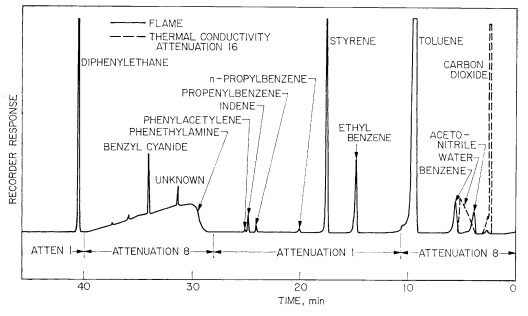


FIGURE. Gas chromatogram of phenylalanine pyrolysate on DC200 silicone oil.

in the pyrolysis of amine to olefin. Each of the products indicated was identified by comparison of its mass spectrum with published spectra.⁴

Pyrolysis of tyrosine under comparable conditions (4 sec. at 500°), gave the expected products phenol, p-cresol, and p-ethylphenol (corresponding to benzene, toluene, and ethylbenzene from phenylalanine). Also formed were the previously reported methane,³ water,³ carbon dioxide,³ and toluene,¹ methyl cyanide, and rearrangement 3-methylindole, and vinylindole. Histidine would be expected to form imidazole and imidazole derivatives on pyrolysis. An established gas chromatographic method⁵ for these was used, but under operating conditions compatible with the mass spectrometer inlet, clearly defined peaks were not obtained. A broad background spectrum of imidazole was obtained (m/e 68, 41, and 40).

(Received, May 23rd, 1968; Com. 654.)

1042

¹C. Merritt, jun. and D. H. Robertson, J. Gas Chromatog., 1967, 5, 96.

² K. Kanomata and Y. Mashiko, J. Chem. Soc. Japan, 1966, 87, 57.

⁸ J. Vollmin, P. Kriemler, I. Omura, J. Seibl, and W. Simon, Microchem. J., 1966, 11, 73.

⁴ ASTM Committee E-14 on Mass Spectrometry, "Index of Mass Spectral Data," American Society for Testing and Materials, Philadelphia, 1963.

⁵ R. Tham, J. Chromatog., 1966, 22, 245.