NOVEL SYNTHESIS OF CHRYSANTHEMUMDICARBOXYLIC ACIDS

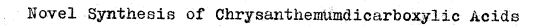
TOSHIO SUGITA

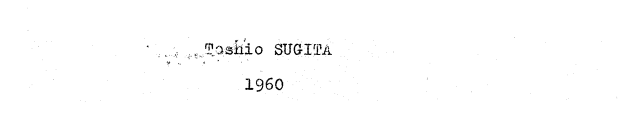
1960

CONTENTS

- I. Introduction
- II. Relative reactivity of the conjugated diene carboxyl compounds
- III. Theory of the addition of aliphatic diazo-compounds to diene system
- IV. Synthesis and stereochemistry of α -methylmuconic acid
- V. Addition of dimethyldiazomethane to methyl \mathfrak{a} -methyl-muconate

- 1 -



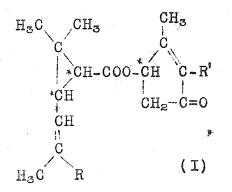


I. INTRODUCTION

It has been well known that the insecticidal constituent of pyrethrum flower, Chrysanthemum cinerariifolium, is a mixture of cyclic keto-esters, referred to as "pyrethrins". The insecticidal activity of "pyrethrins" is so specific in respects of its quick knockdown effect, of the nontoxicity against mammals inspite of the incomparably high insecticidal effect and of producing no resistance in insects. With these advantages it has been practically used as one of the most important insecticides.

The first and comprehensive study on "pyrethrins" was developed by Staudinger and Ruzicka. In 1924 they published their excellent studies¹⁾ on the separation of the active constituents in Dalmatian pyrethrum flowers and the assignment to their structures. Since then, all the expansion in this field, has been based on these works. Although they pioneered the synthetic and structural study on naturally occurring pyrethrins, a quarter century after, their conclusions were partly suffered very important corrections, of course, not that the conclusions were denied wholly. After the Second World War, LaForge and Harper brought a great advance in the structural and synthetic problems of pyrethrins. In consequence, it was established that the naturally occurring pyrethrins consist of a mixture of four substances, i.e. pyrethrin-I, pyrethrin-II, cinerin-I and cinerin-II. It is currently recognized and the structures of these constituents were determined as follows.

- 2 -

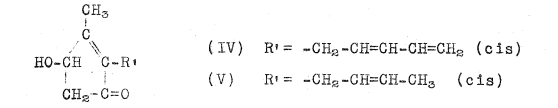


Pyrethrin-I(I) $R = -CH_3$ $R' = -CH_2-CH=CH-CH=CH_2$ (cis)Pyrethrin-II(I) $R = -COOCH_3$ $R' = -CH_2-CH=CH-CH=CH_2$ (cis)Cinerin-I(I) $R = -CH_3$ $R' = -CH_2-CH=CH-CH_3$ (cis)Cinerin-II(I) $R = -CH_3$ $R' = -CH_2-CH=CH-CH_3$ (cis)

These four structures have some common features, that is,all of them are esters, their acid fragments are the monoterpenic acids with cyclopropane nucleus, chrysanthemummonocarboxylic acid (chrysanthemic acid, II) and chrysanthemumdicarboxylic acid monomethylester (pyrethric acid, III'), and their alcohol moieties are cyclopentenolone derivatives, pyrethrolone (IV) and cinerolone (V). These fragments except one had already been synthesized and their geometrical configurations had also been confirmed. The remaining fragment, chrysanthemumdicarboxylic acid, was recentry synthesized by Inouye²⁾ and also by Harper³.

 $\begin{array}{cccc} CH_{3} & & CH-CH=C < R \\ CH_{3} & & CH-CH=C < R \\ & & CH \\ & & COOH \end{array} \quad (II) \quad R = -CH_{3} \quad (trans about the cyclopropane ring) \\ & & (III) \quad R = -COOH \quad (""") \\ & & (III) \quad R = -COOCH_{3} \quad ("") \end{array}$

- 3 -



Chrysanthemumdicarboxylic acid (III) was at first isolated from Japanese pyrethrum flowers by Fujitani⁴⁾ as early as in 1909, and the melting point (164⁰) thereof was reported, but no further study was pursued. The structure of this crystalline dextrorotatory dicarboxylic acid was established by Staudinger and Ruzicka¹⁾ as (+)-trans-3-(2'carboxyprop-l'-enyl)-2:2-dimethylcyclopropane-l-carboxylic acid by ozonolysis to give (-)-trans-caronic acid and pyru-On cold sodium methylate fission of the crude vic acid. semicarbazone of pyrethrins they isolated the half methyl ester of chrysanthemumdicarboxylic acid (pyrethric acid), from which methyl pyruvate was obtained as well as (-)trans-caronic acid on ozonolysis. Therefore it was decided that the cyclopentenolone fragment was esterified with the carboxyl group directly attached to cyclopropane ring, namely the structure III' was established for pyrethric acid.

S. C. L.

ASS.

1407.6

e enter Care e la

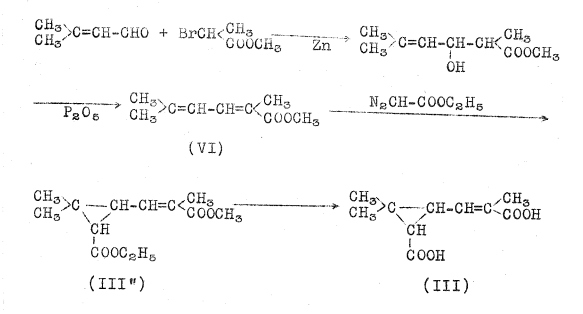
n Ann. Ior anns

61. L.)

Afterwards, Harper^{3,5)} and Inouye⁶⁾ established the absolute configuration of the naturally occurring (+)-transchrysanthemumdicarboxylic acid (III) on the configurative correlation for (+)-trans-chrysanthemic acid (II). But the stereochemistry of the side chain double bond was left unsettled.

The total synthesis of chrysanthemumdicarboxylic acid was recently succeeded by Inouye²⁾ and Harper³⁾ almost concurrently but independently, pursuing the following scheme. Afterwards, Matsui⁷⁾ also obtained the isomeric acids by the other process.

- 4 --

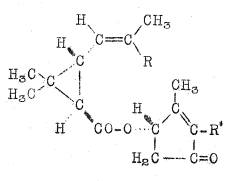


The authors^{8,9)} established the geometrical configuration of the side chain double bond to be trans by the synthetic procedures. The geometrical configuration of $\alpha\beta$ -double bond of $\alpha\delta$ -dimethylsorbic acid and its ester (VI) was decided to be trans by physico-chemical and chemical evidences. As well, this configuration was maintained to chrysanthemumdicarboxylic acid (III).

Also recentry Katsuda¹⁰⁾ decided the absolute configuration of pyrethrolone (IV) and cinerolone (V). Thus the final and complete structures of the naturally occurring pyrethrins were established as follows.

rffester

e. aa



- 5 -

As the results of these developements, the chemical structures and configurations of the constituents of pyrethrum flowers have already been clarified and their total syntheses have further been accomplished. Although chrysanthemumdicarboxylic acid have four geometrical isomers theoretically, Inou/e and Harper have obtained two of them, i.e. trans, trans- and cis, trans-isomers* which are the whole available from their synthetic procedures since the trans-isomer of a*-dimethylsorbic acid alone has been obtainable as the starting material of the synthesis. Therefore, some other procedure has been required to obtain the remaining isomers. Beside this, the above-mentioned method is a complicated one, and there yet remains room for improvement in the yield.

4

二 《本記書 世

注意構成が

3522333

L'ENR

The authors¹¹⁾ succeeded in the synthesis of trans, trans- and trans, cis-chrysanthemumdicarboxylic acids by a novel and stereospecific route of synthesis and established their geometrical configurations, which will be described in details in the following chapters. The latter isomer, trans, cis-chrysanthemumdicarboxylic acid could not be synthesized by Inouye's method. It is the further advantage of the novel route of synthesis that the each geometrical isomer of the acid is obtainable without contamination of the other isomers by more facile procedures.

II. RELATIVE REACTIVITY OF THE CONJUGATED DIENE CARBOXYL COMPOUNDS.

The author will be concerned in this chapter with the

* Regarding the representation of configurations, refer to the footnote on page 31.

- 6 -

reactivity of conjugated unsaturated systems, in particular the addition reactions to conjugated diene carboxylic acids. Reactions of the simple unsaturated systems have been kinetically studied, and also those of the simple conjugated diene compounds such as butadiene have been revealed to some extent. However, reactions of the conjugated diene carboxyl systems, especially those of substituted ones present a more complicated problem and have not completely been elucidated as yet.

The addition reactions of symmetrical molecules, such as hydrogen which is considered as an electrophile, and of Diels-Alder type to conjugated dienes have been discussed already in many papers. Regarding the orientation of addition of these types, as is well known, 1,4"addition takes place more commonly. In nucleophilic additions to the conjugated polyene carbonyl systems, the orientation of addition takes place in somewhat different and complicated aspects. The orientation of reductions of conjugated dienoic acid by dissolving metals was more favourable in 1,4-addition. The mechanism of this reductions has been regarded as a nucleophilic one. On reduction with sodium amalgam, muconic acid, the simplest butadiene derivative having terminal carboxyl groups HOOC-CH=CH-CH=CH-COOH, gave predominantly the β Y-unsaturated dihydro-compound.¹²⁾ β -Vinylacrylic acid CH2=CH-CH=CH-COOH also gave ao-dihydro-derivative by the same reducing agent in alkali medium, however, its methyl derivatives gave $\alpha\beta$ - as well as $\alpha\delta$ -adducts on the reduction, and the proportion of $\alpha\beta$ - to $\alpha\delta$ -adducts varied with the position of the substituent.¹³⁾ On the other hand, additions of Michael type, for example, of a malonic or cyanoacetic ester to sorbic ester, are oriented to either the β - or the δ -position,¹⁴⁾ since these positions are in principle positively polarisable, and are therefore potential positions of

たい対応

2.50000

网络美国语

3.13 P.1

X WY De

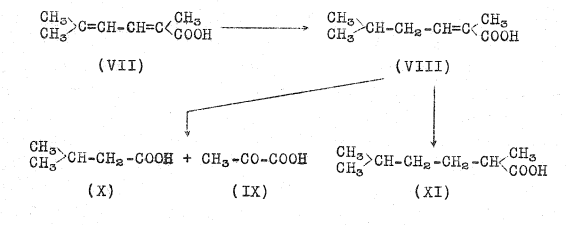
-- 7 --

attachment of the anionic portion of the addendum.

As mentioned above, the relative reactivity of conjugated polyene system is very complicated and undergoes a change by the substituents. Therefore, the author had to deal with some reactions in relation to these problems.

§ Semihydrogenation of $\alpha\delta$ -dimethylsorbic acid.⁸⁾

At first, the partial hydrogenation of $\alpha\delta$ -dimethylsorbic acid (VII) to give $\alpha\delta$ -dimethyl- $\Delta\alpha$ -hexenoic acid (VIII) is described. Farmer and Hughes¹⁵⁾ reported that sorbic acid, when submitted to 50% hydrogenation in the presence of palladium catalyst, gave a mixture of dihydroacids in which the $\Delta\alpha$ -dihydrosorbic acid predominated ranging from 85 to 90%, whilst with a platinum catalyst only a mixture of an almost equal amount of fully reduced and unchanged conjugated compounds was obtained, the production of small proportions (below 20%) of dihydro-compounds being recorded.



r Lebendiji V letense

the start

j. . . . ? .

In the present author's case, the semihydrogenation of $a\delta$ -dimethylsorbic acid (VII) over Pd-BaSO₄ catalyst predominantly afforded the Aa-dihydro-compound, that is $\gamma\delta$ -addition was favoured. $a\delta$ -Dimethyl-Aa-hexenoic acid (VIII) was obtained in the yield of 73% after repeated purifications by

means of partial esterification * and rectification. The product was characterized by ozonolysis to yield pyruvic acid (IX) and iso-valeric acid (X). In the UV-spectrum of the resulting α^{3} -dimethyl- $\Delta \alpha$ -hexenoic acid, the occurrence of the single intense band (Amax. 218 mu, & 14,300), characteristic for the $\alpha\beta$ -unsaturated carboxylic chromophore, together with the disappearance of the conjugated diene carboxylic band (λ max. 273 m μ) which was exhibited by the parent dimethylsorbic acid (VII), were consistent with the chemical evidence mentioned above. The fact that the extinction coefficient at this band did not alter on further purification. excluded the contamination of the possible $\Delta\beta$ -, $\Delta\gamma$ -dihydroor fully reduced compounds, which were spectrally inert in : this region. This acid was quantitatively hydrogenated over a platinum catalyst and was shown to absorb one equivalent hydrogen, yielding the known $a\delta$ -dimethylcaproic acid (XI)¹⁶⁾

teres to see all the second se

1.753

in fer e

430

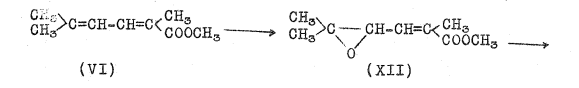
The equivalent weight and the acid dissociation constant were also determined.⁸⁾ All criteria supported purity of this acid.

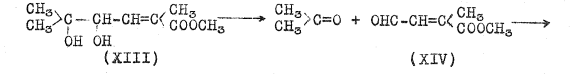
§ Selective oxidation of αδ-dimethylsorbic acid⁹ Heinänen¹⁷ reported that the selective epoxidation occured at the Yδ-double bond of methyl sorbate by the action of perbenzoic acid in the cold, giving methyl Y8-epoxy-Δαhexenoate.

The epoxidation of our methyl αδ-dimethylsorbate (VI) by perbenzoic acid gave methyl Υδ-epoxy-αδ-dimethyl-Δα-

* In the sense that $\Delta\beta$ - and $\Delta\gamma$ -dimethylhexenoic and dimethylcaproic acids become esterified, whilst $\Delta\alpha$ -dimethylhexenoic acid remains unchanged. (Compare Ecott and Linstead's modification of Sudborough's method, J.Chem.Soc., <u>1929</u>,2153; <u>1932</u>,125)

- 9 -







-1897 - S.H.

Sa ^enA

ېلې د مېښې د د د د د د

hexenoate (XII) as was expected, though this addition seems to proceed by a free-radical mechanism, one atom of oxygen being taken up at the Yo-double bond of the parent ester. The Yo-epoxy structure of the product can reasonably deduced from the subsequent degradations. Conversion of the epoxy-ester into the dihydroxy-compound was easily effected by the treatment with diluted sulphuric acid, yielding $\gamma \delta$ dihydroxy-a6-dimethyl-1a-hexenoate (XIII). Lead tetraacetate cleaved the glycolic carbon-carbon linkage of XIII and gave (α) -methyl mesaconaldehydate (XIV), which was characterized by 2,4-dinitrophenylhydrazone, mp. $204-4.5^{\circ}$. (a)-Methyl mesaconaldehydate was then oxidized with peracetic acid to (α)-methyl (β)-hydrogen mesaconate (XV), mp. 82-3⁰, which as well as the amide, mp. 117°, derived therefrom, were completely consistent with the literature¹⁸⁾ Cold saponification of (α) -methyl (β) -hydrogen mesaconate gave mesaconic acid (XVI), mp. 202-3°, not depressed by mixed melting point comparison with an authentic specimen.¹⁹⁾

- 10 -

The direct peroxidation of $\alpha\delta$ -dimethylsorbic acid (VII) by means of hydrogen peroxide, also gave mesaconic acid (XVI) but in an inferior yield. This result indicated that the selective oxidation took place at the same $\gamma\delta$ -double bond as in its epoxidation.

§ Addition of ethyl diazoacetate to $\alpha\delta$ -dimethylsor.

In the system of $\alpha\delta$ -dimethylsorbic acid (VII), as mentioned above, the Y\delta-double bond is more reactive than is the the $\alpha\beta$ -double bond, and this result is supported also by the fact that ethyl diazoacetate adds exclusively to Y\delta-double bond of $\alpha\delta$ -dimethylsorbate (VI), yielding chrysanthemumdicarboxylate (III"). Discussions on the mechanism of this reaction will be related in the next chapter.

§ Lactonization of α -methylmuconic acid²⁰.

When α -methyl-cis,cis-muconic acid (XVII c-c) was treated with cold 80% sulphuric acid or boiled with water, an unsaturated lactonic acid, Y-carboxymethyl- α -methyl- $\beta\alpha$ butenolide (XVIII) alone was obtained. Experiments and its structural arguments will be described in chapter III. Although two kinds of lactonic acids are to be expected for this acid, namely the lactonic acid in which α -carboxyl group is concerned in the lactonization and the other in which δ -carboxyl group is concerned, only one of which has exclusively been obtained.

 $HOOC-CH=CH-CH=C < CH_3 COOH$

 $HOOC-CH_{z}-CH \langle CH=C-CH_{z}$

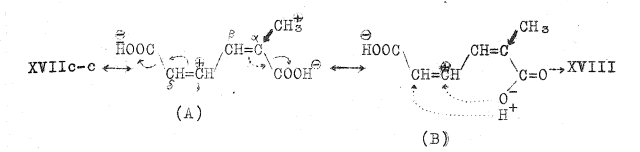
(XVII c-c)

(XVIII)

The uniformity of the lactonization of the cis, cis-acid

- 11 -

can be interpleted as follows. In the starting acid (XVII c-c), the electrons are oriented as indicated in (A). The



electron attractive effect of the carboxyl group, attached to the δ -carbon atom, induces the polarization of the adjascent double bond producing a positive charge on the γ -carbon atom. The same effect of the carboxyl group attached to the a-carbon atom, also produces a positive charge on the β carbon atom, which is much more weakened by the electron releasing effect of the a-methyl substituent, and hence, the positive charge on the β -carbon atom is weaker than that on the γ -carbon atom. Consequently, the attack of the polarized a-carboxyl oxygen to the γ -carbon atom is preferred and then followed by the addition of proton to the δ -carbon atom, resulting in the formation of the only known lactonic acid (XVIII).

From the fact that the $\Upsilon\delta$ -double bond was more reactive than the $\alpha\beta$ -double bond in α -methylmuconic acid system as mentioned above, it was resonably expected that the addition of dimethyldiazomethane to the $\Upsilon\delta$ -double bond of α -methyl- α muconic acid should be favoured. Regarding this problem, the author will discuss in the next chapter in details. Indeed, this was the case with the addition reaction, thus verifying the proposed mechanism as well as the preferred reactivity of the $\Upsilon\delta$ -ethylenic linkage in this system.

in the second of

w chif

10045

- 12 -

EXPERIMENTAL

Semihydrogenation of $\alpha\delta$ -dimethylsorbic acid.

A 15.0 g quantity of $\alpha\delta$ -dimethylsorbic acid (VII; mp. 134-5°)^{2a)} dissolved in 150 ml of methanol was hydrogenated over Pd-BaSO₄ catalyst (1.2 g) in a shaking apparatus until 2564 ml (at 19°) of hydrogen (equivalent to one double bond) was absorbed. The reduction product thus obtained, was freed from the catalyst and solvent, and carefully distilled under reduced pressure. The distillate was submitted to partial esterifications in order to eliminate the possible $\Lambda\beta$ - and $\Delta\gamma$ -dihydro acids; the distillate (14.3 g, 95%) was mixed with 0.2 N-ethanolic hydrogen chloride (70 ml) and kept at room temperature for 5.5 hrs. After the duration, the solution was diluted with four times its bulk of water, made faintly alkaline against lithmus paper by the addition of sodium carbonate. The neutral substance separated was collected with ether, then the aqueous layer was concentrated under a reduced pressure below 50° until free from alcohol. The neutral fraction was again completely removed with ether at this stage. Acidification of the aqueous residue and thorough extraction with ether, followed by drying and removal of the solvent, gave the fraction bp. 118-1220/10 mm (11.1 g, 73%). This crop was enough pure at this stage, but in order to obtain the sample of the highest purity for physico-chemical measurements which were needed for determination of the geometrical configuration of this acid, this was again subjected to partial esterification exactly in the same manner as described above and in further rectifications,

yat da ki

. Alatha

odýme.

* For physico-chemical determination of the geometry, which may be applicable to the conjugated diene carboxylic acid in general, see the detailed article by the same authors⁸)

- 13 -

only a center cut of pure $\alpha\delta$ -dimethyl- $\Delta\alpha$ -hexenoic acid was collected, bp. 120.5-121.5°/10 mm (6.3 g); n_D²⁵ 1.4597; equivalent weight: Found 141.1, Calcd. for $C_7H_{13}COOH$ 142.2; $\lambda \max$. 218 m/A, ε 14,300. It crystallized in a prism when chilled in dry-ice and melted at about 0° . p-Phenylphenacylester, mp. 54-6° (Anal. Found C 78.55, H 7.13, Calcd. for C22H2403 C 78.54, H 7.19). Distillation of the neutral ether extracts combined gave the fraction of ester, bp. $84-88^{\circ}/22$ mm; n_{D}^{25} 1.4313 (4.5 g). It decolourized the permanganate solution at room temperature and absorbed bromine instantly. However, this was not subjected to further investigation. Ozonization: --- One gram of the dihydro-acid in 50 ml of chloroform was treated with an excess of ozone at 0° . The solvent was removed in vacuo, and the remaining ozonide was decomposed with water on a water bath for 10-15 mins. To this aqueous solution, was then added the solution of 2,4dinitrophenylhydrazine in dilute hydrochloric acid to precipitate any carbonyl compound present, and was kept overnight. The yellow precipitate was collected and recrystallized several times from ethanol to give 2,4-dinitrophenylhydrazone of pyruvic acid, mp. 218° (0.6 g). The melting point was not depressed by admixture with an authentic specimen. The filtrate from the hydrazone was thoroughly extracted with ether and the ether solution was dried over anhydrous sodium sulphate, and after removal of ether, the residue was distilled to give isovaleric acid (0.3 g), bp. 170-175°; n_{D}^{22} 1.4020. p-Phenylphenacylester, mp. 77-8° (cf. Drake²¹⁾ mp. 76°). The melting point was not depressed by admixture with an authentic specimen.

and the second

C. P. S.

*新,我*没有。

10 march

Car Leve

北京工業協力

计分表类总统

动的标志。

. Con

steine)

t sina

Quantitative hydrogenation: A 0.233 g quantity of the acid (bp. $120.5-121.5^{\circ}/10$ mm) in 30 ml of ethanol was hydrogenated over platinum oxide catalyst (11 mg) in a shaking hydrogenation apparatus, and absorbed 40.0 ml (at 27°) of

- 14 -

i. Ísti AGEL D 100 (20)£ ار ایک ایک در ایک این میں میں کہ آج ای 131.15、新闻日 and the first and 言い現象が、 1.0788. 心正盘 s (Cé)VQ AT LY 法法法法法 发。 Lasse Carre

hydrogen, equivalent to one double bond. The reduction product was freed from both catalyst and solvent, and distilled to give the fully reduced $\alpha\delta$ -dimethylcaproic acid (XI)¹⁶) in a yield almost quantitative, bp. 115-116°/13 mm; n_D²⁰ 1.4261. Amide, mp. 102-3°; p-phenylphenacylester, mp. 66°, identified by mixed melting point comparison with authentic specimens, respectively.

Selective oxidation of $\alpha\delta$ -dimethylsorbic acid.

Methyl Yô-epoxy- α ô-dimethyl- $\Delta \alpha$ -hexenoate (XII) Four grams of methyl α ô-dimethylsorbate (VI; 0.027 mole.)^{2a)} were dissolved in 5 ml of dry chloroform and to this were added 147 ml of 3.5% perbenzoic acid in chloroform (0.038 mole.). The mixture was kept at 0° and the consumption of the active oxygen was estimated by means of iodometry on a small portion drawn from the reaction mixture at intervals. After 5 d days' standing, 0.03 atom of axygen was taken up, then the excess of perbenzoic acid was decomposed with sodium sulphite, removed by washing with sodium carbonate and dried over anhydrous sodium sulphate. After removal of the solvent, the residue was distilled under a reduced pressure to give methyl Yô-epoxy- α ô-dimethyl- $\Delta \alpha$ -hexenoate, bp. 91-2°/6 mm; n_D^{20} 1.4672; yield 3.7 g (84%).

Methyl Yδ-dihydroxy-αδ-dimethyl-Δα-hexenoate (XIII) To a 1.5 g quantity of epoxy-ester (XII), was added 0.5 ml of 5% sulphuric acid, kept at room temperature and after several hours it turned homogeneous. This was then extended in 100 ml.of alcohol free ether and was completely dried over anhydrous sodium sulphate. Removal of ether gave methyl Yδ-dihydroxy-αδ-dimethyl-Δα-hexenoate in quantitative yield. Plates (from methanol and benzene); mp. 50-1°; Anal. Found C 57.40, H 8.46, Calcd. for C₉H₁₆O₄ C 57.43, H 8.57. (α)-Methyl mesaconaldehydate (XIV) The dihydroxy-

- 15 -

ester (XIII; 1.48 g, 0.0079 mole.) was dissolved in 100 ml of dry benzene and to this were added 3.9 g of freshly prepared lead tetraacetate (0.0087 mole.) in three portions. The mixture was stirred at 50° for 2 hrs. and then at 60° for 2 hrs. After the duration, excess of lead tetraacetate was decomposed with water and lead oxide formed was removed by filtration. The filtrate was dried over anhydrous sodium sulphate and the solvent was distilled off. Distillation of the residue in vacuo gave (α)-methyl mesaconaldehydate, bp. 76-8°/12 mm; n_D^{20} 1.4680 (0.75 g, 75%). 2,4-Dinitrophenyl-hydrazone, orange needles (from methanol), mp. 204-4.5° (decomp.), (Anal. Found C 46.73, H 3.92, N 18.21, Calcd. for $C_{12}H_{12}O_6N_4$ C 46.76, H 3.92, N 18.18).

(a)-Methyl (β)-hydrogen mesaconate (XV) The aldehydeester (XIV; 0.3 g, 0.0024 mole.) was dissolved in 2.7 ml of 13.3% peracetic acid (0.0047)mole.) and the mixture was kept cold for 36 hrs. Then the mixture was diluted with water and dried up in vacuo to yield a crystalline mass. The residue was crystallized from petroleum ether (bp. 40-50°) to give (α)-methyl (β)-hydrogen mesaconate (0.31 g), needles, mp. 82-3°. Anal. Found C 50.04, H 5.61, Calcd. for C₆H₈O₄ C 50.00, H 5.60. Amide, needles (from ether-petroleum ether), mp. 115.5-6.5° (Anal. Found C 50.39, H 6.51, N 9.72, Calcd. for C₆H₉O₃N C 50.34, H 6.34, N 9.79). These were perfectly consistent with the literature.¹⁹

Mesaconic acid (XVI) (α)-Methyl (β)-hydrogen mesaconate (XV; 0.1 g) was hydrolysed with cold 5% ethanolic potassium hydroxide, dried up in vacuo and acidified with dilute sulphuric acid. This was extended in ether, dried over anhydrous sodium sulphate and the removal of ether gave mesaconic acid, cubes (from water), mp. 202-3°. Anal. Found C 46.07, H 4.60, Calcd. for C₅H₆O₄ C 46.16, H 4.65. The melting point was not depressed by **the mixed** melting point-

- 16 -

comparison with an authentic specimen prepared by the recorded method.¹⁹⁾ bis-p-Phenylphenacylester, fine needles (from chloroform), mp. 204-5°. (Anal. Found C 76.35, H 5.14 Calcd. for C₃₃H₂₆O₆ C 76.43, H 5.05).

Peroxidation of a0-dimethylsorbic acid (VII) a0-Dimethylsorbic acid (VII; 0.5 g) was dissolved in 10 ml of dilute acetic acid, and to this were added 5 ml of 30% hydrogen peroxide. The mixture was warmed on a steam bath for ca. 10 hrs. with additional peroxide at intervals. At the end of this time, the solution was dried up in vacuo and the solid residue was extracted with cold ether. To the remaining residue were added a drop of dilute sulphuric acid and 2 ml of water, and it was warmed on a water bath. Reduction of the volume gave mesaconic acid (19 mg), which melted at 201-3° after recrystalization from water. The melting point was not depressed by admixture with an authentic specimen. Further reduction of the filtrate gave a less pure product (ca. 25 mg).

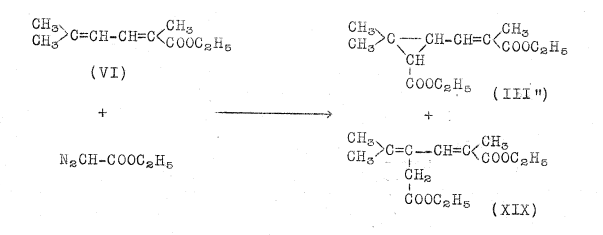
III. THEORY OF THE ADDITION OF ALIPHATIC DIAZO-COMPOUNDS TO DIENE SYSTEM.

The mechanism of the addition of almphatic diazo-compounds to olefins remains uncertain. It is currently interpreted by two ways, i.e. the participation of the free radicals and the path over pyrazoline intermediates, and the former seems to be used more widely for the interpretations at the present.

Inouye²²⁾ isolated a pyrazoline intermediate on the addition of diazoacetate to $\alpha\delta$ -dimethylsorbate (VI), then he preferred the pyrazoline intermediate mechanism for that

- 17 -

addition reaction. He also took the fact for the explanation that the addition of ethyl diazoacetate to ethyl $\alpha\delta$ -dimethyl-sorbate (VI) produced an acyclic isomer (XIX) as well as the cyclopropane derivative, i.e. ethyl chrysanthemumdicar-boxylate (III").

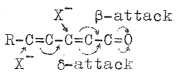


The formation of the acyclic isomer (XIX) could not be explained by means of the addition of the diradical :CHCOOC₂H₅. By assuming the pyrazoline intermediate (XX), the formation of the all addition products could be satisfactorily explained.

As mentioned in chapter II, Inouye^{2a)} obtained only the Yô-adduct in the addition of ethyl diazoacetate to ethyl aôdimethylsorbate. aô-Dimethylsorbic acid (VII) also indicated the more reactivity at Y8-double bond in the catalytic hydrogenation and oxidations. Therefore, in order to explain the exclusive addition of diazo-compound to the Yôdouble bond, the following mechanism seems appropriate. Either the β - or the δ -position in the conjugated compounds such as β -vinylacrylic acid and sorbic acid, is in principle positively polarisable,'and is, therefore, a potential place of attachment of the anionic portion of the addendum, especially the δ -position is more potential in the substituted

- 18 -

sorbic acid systems,



Whilst, diazoacetate takes part in the reaction under consideration as the resonance hybrid,

- + N=N-CH-COOC₂H₅

Thus, in due course, the anionic portion of diazoacetate enters the δ -position, and then the protonic part becomes bound at the Y-carbon atom. As the result of the addition of these two components, there forms the pyrazeline (XX), which is responsible for producing both ethyl chrysanthemumdicarboxylate (III") and its acyclic isomer (XIX). This

mechanistic argument can apply to the reactions, such as the addition of ethyl diazoacetate to methyl $\alpha\beta\delta$ -trimethylsora. bate²³⁾ and methyl sorbate²⁴⁾ and of diphenyldiazomethane to menthyl sorbate.²⁵⁾ All of these olefinic components had the same conjugated system as mentioned above, and each diazo-compound added exclusively to the $\gamma\delta$ -double bond. Addition

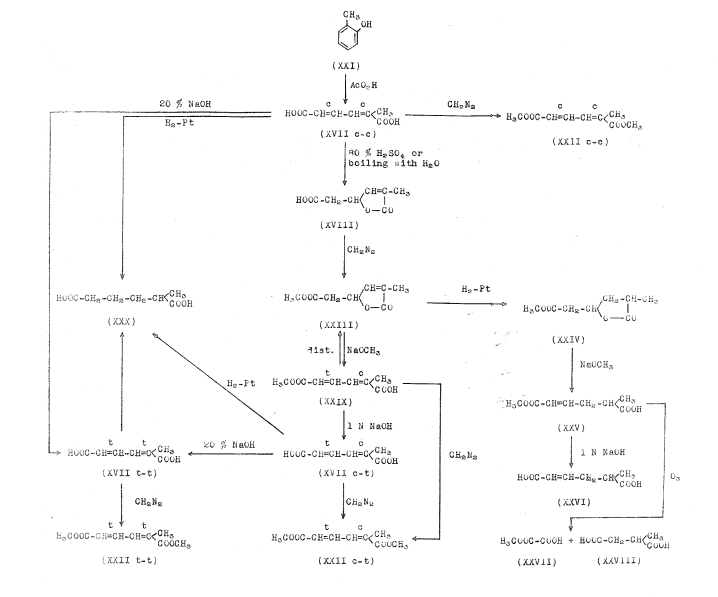
- 19 -

of ethyl diazoacetate also took place at Y8-ethylenic bond of 1-phenylbutadiene 26 which had an similar conjugated system as sorbic acid. In contrast, Guha and Sankaran²⁷⁾ observed terminal addition of diazomethane to ethyl muconate. However, two ethylenic bonds of ethyl muconate are equally poralized and there is no difference of the reactivity in each double bond, therefore the result seems to be reasonably expected. Should this theory be applied to the system of α -methylmuconate, in which the YS-double bond is more reactive than $\alpha\beta$ -double bond as mentioned in the preceding chapter, diazo-compound should add to the Y8-double bond of a-methylmuconate. Hence it must become possible to obtain chrysanthemumdicarboxylic acid by means of the addition of dimethyldiazomethane to a-methylmuconate. In the fact, the addition reaction did take place at the Yo-double bond. giving chrysanthemumdicarboxylic acid. The details of this reaction will be described in chapter V.

IV. SYNTHESIS AND STEREOCHEMISTRY OF α -METHYL-MUCONIC ACID²⁰

The theoretical consideration mentioned in the preceding chapters leads to a result that it is possible to synthesize isomeric chrysanthemumdicarboxylic acid by the addition of dimethyldiazomethane to a-methylmuconate. Hence, it became necessary to prepare a considerable amount of amethylmuconic acid as a starting material, and to establish the geometrical configuration thereof for furthering this scheme. Kuhn and Michel²⁸⁾ prepared an isomer of a-methylmuconic acid by the condensation of ethyl tiglate with ethyl oxlate, followed by acetylation and hydrolysis. The geome-

- 20 -



trical configuration of the acid, however, has not yet been confirmed, though it is assumed tentatively to be trans, trans.

a-Methylmuconic acid can exist theoretically in four geometrically isomeric forms cis,cis; cis,trans; trans,cis and trans,trans. Of these, three isomers were obtained by the present author through a novel route of synthesis and their geometrical configurations were established. The isolation of the remaining one isomer, trans,cis, has failed in spite of any attempt.

The oxidation of o-cresol (XXI) with peracetic acid gave α -methyl-cis,cis-muconic acid (XVII c-c), mp. 189-90°, which was converted with diazomethane into the dimethylester (XXII c-c), mp. 35°.

The treatment of the cis, cis-acid (XVII c-c) with cold 80% sulphuric acid gave the unsaturated lactonic acid (XVIII), mp. $105-6^{\circ}$, the structure of which is evidenced by the subsequent chemical processes. The lactonization was also effected by boiling with water, yielding the same compound (XVIII). It was of interest that the lactonization of the acid (XVII c-c) under any reaction conditions employed, gave exclusively the lactonic acid (XVIII) alone. This seems very likely, from the theoretical consideration which was discussed in chapter II.

.

The lactonic acid (XVIII) afforded a liquid methyl ester

 * The nomenclature of these compounds are based on αtrans cis
 methyl-muconic acid, therefore, e.g. MeOOC-CH=CH-CH=CMe-COOH
 is named (δ)-methyl (α)-hydrogen α-methyl-cis,trans-muconate.
 The terms cis and trans are given in positional order, thence
 the first term indicated the configuration of the αβ-double
 bond.

- 21 -

(XXIII) with diazomethane. The unsaturated lactonic ester (XXIII) took up one mole. hydrogen on catalytic hydrogenation, giving the saturated lactonic ester (XXIV), mp. $45-6^{\circ}$. The treatment of the saturated lactonic ester (XXIV) with sodium methoxide in methanol gave an acyclic isomer (XXV), hydrolysis of which afforded a-methyl- Δ Y-dihydromuconic acid (XXVI), mp. 121-2°. The half ester (XXV) was characterized by ozonolysis to yield methyl hydrogen oxalate (XXVII) and methylsuccinic acid (XXVIII). Therefore, the structures of the unsaturated lactonic acid and of the compounds derived therefrom, were confirmed as XVIII - XXVI.

The treatment of the unsaturated lactonic ester (XXIII) with sodium methoxide in methanol gave the expected (δ)methyl (α)-hydrogen α -methyl-cis,trans-muconate (XXIX), mp. 121°, which was readily reconverted into the parent unsaturated lactonic ester (XXIII) by distillation under reduced pressure. On hydrolysis with dilute sodium hydroxide, the half-ester (XXIX) produced α -methyl-cis,trans-muconic acid (XVII c-t), mp. 172°. Esterification of either the acid (XVII c-t) or the half-ester (XXIX) with diazomethane yielded the same dimethyl ester (XXII c-t), mp. 60°.

When the cis,cis- (XVII c-c) and the cis,trans-acids (XVII c-t) were heated with 20% sodium hydroxide, these were equally isomerized into the trans,trans-form (XVII t-t), mp. 273°, which might be identical with that obtained by Kuhn and Michel²⁸, mp- 276°, though direct comparison by mixed melting point is lacking. The acid (XVII t-t) was converted with diazomethane into the dimethyl ester (XXII t-t), mp. 55.5°, which also might correspond to the ester recorded by Harris and Binns²⁹, mp. 46-7°.

On catalytic hydrogenation over platinum catalyst, each of the trans, trans-, cis, trans- and cis, cis-acids took up two moles hydrogen and gave the same α -methyladipic acid

- 22 -

(XXX), mp. $58-9^{\circ}$, which was also characterized by its diamide, mp. $185-6^{\circ}$.

Configuration: — The configurations of the three acids, (XVII c-c), (XVII c-t) and (XVII t-t), followed unambiguously from the method of preparation and the spectral data. The high melting and the most stable isomer (XVII t-t) is expected to have all trans-configuration, because it is also obtained by the condensation of tiglate of the well-defined trans-configuration with oxalate, in which the trans-configuration is favoured in general. The failure of the isomer to be lactonized with 80% sulphuric acid into lactonic acid, is in good agreement with this deduction.

The acid (XVII c-c) obtained by the oxidative ringfission of o-cresol is most likely to have cis,cis-configuration. It is, in fact, readily lactonized to XVIII. Moreover, it is isomerized to the high melting acid (XVII t-t) both directly and via the third acid (XVII c-t).

The half-ester of the third acid (XXIX) produced by ring opening reaction with alkoxide is expected to have cis, trans-configuration, The $\alpha\beta$ -double bond must be undoutedly cis in the lactone. The cis-configuration of the $\alpha\beta$ -double bond should be retained as such during the ring opening reaction, since the resulting half-ester was capable of ready lactonization into the parent compound (XXIII). The Yô-double bond which is created by the ring opening reaction, is considered to have trans-configuration by the analogies in similar reactions.

The third acid (XVII c-t) can also be isomerized to the trans, trans-acid (XVII t-t). It is evident, therefore, that the third acid (XVII c-t) has cis, trans-configuration.

The first acid (XVII c-c) produced by ring fission of o-cresol has at least one cis-double bond, since it is readily lactonizable. It can be isomerized to the cis, trans-

isomer via the lactone, hence the trans, cis-configuration is excluded for it. Therefore, the first acid (XVII c-c) should reasonably be concluded to be cis, cis, also by the expectation that it is formed by the fission of an aromatic ring under a mild condition, then it will appear in a coiled phase with orientation of cis, s-cis, cis at the first step.

The conclusions on the configurations of each acid are in complete agreement with spectral data.

The UV-spectrum data of the isomeric α -methylmuconic acids and their esters are collected in Table 1, together with those of some isomeric homologues. Positions of the absorption maximum of each isomer are practically the same.

Table 1. Ultra-violet light absorption of isomeric α -methylmuconic acids in comparison with some homologues.

	cis,cis	cis,trans	trans, trans
	λmax. (mμ) ε	λmax. (mμ) ε	Amax. (mµ) E
α -Methylmuconic acid	269 20,700	269 25,700	273 29,700
methylester	270 24,800	272.27,600	275 31,100
Muconic acid ³⁰⁾	258 17,000	259 25,600	259 29,100
methyl ester	259 26,400	260 29,800	259 36,700
acid ³¹⁾	281 20,400	280 25,500	282 31,450
methyl ester	280 24,750	280 31,600	282 33 ,300
β -Methylmuconic acid ³²⁾	utay ¹ , date	265 19,000	265 22,400
methyl ester		265 22,100	266 28,400

There is no significant cis-shift, but the extinction coefficients show a significant difference in every case,

- 24 -

the coefficient of cis, cis-isomer is the lowest and each cis \rightarrow trans inversion increases the intensity. The validity of these criteria was also discussed in the series of α -methyl- $\alpha\beta$ -unsaturated acids, reported in the previous paper of the authors⁸.

The cis,cis-acid has a maximum extinction coefficient of 20,700 at 269 mµ, the cis,trans-acid, 25,700 at 269 mµ, and the trans,trans-acid, 29,700 at 273 mµ. The methyl esters have the values of 24,800 at 270 mµ, 27,600 at 272 mµ and 31,100 at 275 mµ, respectively. These values unequivocally agreed with the expected values for each configuration.

In Table 2 are summarized the characteristic bands in IR-spectra of the isomers.

Table 2. Characteristic bands in the infra-red absorption spectra of the isomeric α -methylmuconic acids. (cm⁻¹)

cis-cis	cis-trans	trans-trans
	995	988
839	822	820
736	- 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997	

EXPERIMENTAL

a-Methyl-cis,cis-muconic acid (XVII c-c) A mixture of 73.5 g (0.68 mole.) of o-cresol (XXI) and 790 ml of 13% peracetic acid solution (1.36 mole. of peracetic acid) was cooled in ice for 24 hrs. and then kept cold for 1 or 2 weeks. After the duration, the crystalline precipitate was

- 25 -

collected, washed with acetic acid and recrystallized from methanol by cooling with dry-ice, to give the pure a-methylcis,cis-muconic acid in prisms, mp. $189-90^{\circ}$ (this mp. was taken from a bath at 160° , with the temperature rising 10° per minute, since it varied with the rate of heating); λ max. 269 mµ, ϵ 20,700. Anal. Found C 53.73, H 5.26, Calcd. for $C_7H_8O_4$ C 53.84, H 5.16. yield 12.3 g (11.3%)

Quantitative hydrogenation:--- A 0.103 g quantity of the acid (XVII c-c) in 60 ml of methanol was hydrogenated over a platinum oxide catalyst (10 mg) in a shaking apparatus and absorbed 36 ml (at 20°) of hydrogen, equivalent to two double bonds. The reduction product was freed from the catalyst and the solvent, and recrystallized from benzene to give a-methyladipic acid (XXX), mp. 58-9° (Mazza and DiMase³³⁾ recorded mp. 61°). Anal. Found C 52.57, H 7.45, Calcd. for $C_7H_{12}O_4$ C 52.49, H 7.55. Also characterized by diamide, mp. 185-6° (Bouve ault and Locquin³⁴⁾ recorded mp. 186.5°), (Anal. Found C 53.06, H 8.91, N 17.79, Calcd. for $C_7H_{14}O_2N_2$ C 53.14, H 8.97, N 17.71).

Methyl a-methyl-cis,cis-muconate (XXII c-c) The cis, cis-acid (XVII c-c) was treated with ethereal diazomethane solution in the usual manner and the dimethyl ester was obtained in the yield almost quantitative. After recrystallization from light petroleum (bp. 40-50°) or methanol, it formed prisms, mp. 35°; λ max. 270 m μ , ε 24,800. Anal. Found C 58.70, H 6.80, Calcd. for C₉H₁₂O₄ C 58.69, H 6.57.

 Υ -Carboxymethyl- α -methyl- $\Delta \alpha$ -butenolide (XVIII)

(a) A 3.62 g quantity of the cis,cis-acid (XVII c-c) was shaken with 36 ml of cold 80% sulphuric acid for l hr. After 24 hrs., the reaction mixture was poured onto ice and the bulk of the acid was neutralized with an aqueous ammonia (acid to Congo-red). The solution was thoroughly extracted with ether. After removal of ether, the product was recrystallized from chloroform to yield the lactonic acid (XVIII)

- 26 -

in prisms, mp. $105-6^{\circ}$. Anal. Found C 53.60, H 5.34, Calcd. for $C_7H_8O_4$ C 53.84, H 5.16. yield 3.08 g.

(b) A 0.75 g quantity of the cis,cis-acid (XVII c-c) was heated with water under reflux for 1 hr. The reaction mixture was dried up on a water bath and the residue was recrystallized from ethyl acetate to yield Υ -carboxymethyl-a-methyl-Aa-butenolide, mp. 105°. Anal. Found C 53.88, H 5.03, Calcd. for $C_7H_8O_4$ C 53.84, H 5.16. The mixed mp. with the product in the procedure (a) was 105° , yield 0.60 g.

Y-Carbomethoxymethyl- α -methyl- $\beta\alpha$ -butenolide (XXIII) To the lactonic acid (XVIII; 9.21 g, 0.059 mole.) in ether was added an ethereal solution of diazomethane (0.065 mole.) under cooling and well stirring, the product then distilled under reduced pressure to give the lactonic ester, bp. 150-1°/9 mm; n_D^{20} 1.4741. (9.28 g)

Y-Carbomethoxymethyl-a-methylbutanolide (XXIV) A 1.314 g quantity (0.0077 mole.) of the unsaturated lactonic ester (XXIII) in 50 ml of methanol was hydrogenated over platinum catalyst and it took up 218 ml (at 20°) of hydrogen, equivalent to 1.1 mole. The reduction product was freed from the catalyst and the solvent, and distilled to give the saturated lactonic ester, bp. $136-9^{\circ}/5$ mm; it crystallized from ether-light petroleum in laths, mp. $45-6^{\circ}$. Anal. Found C 55.73, H 6.98, Calcd. for $C_8H_{12}O_4$ C 55.80, H 7.03.

 (δ) -Methyl (α) -hydrogen α -methyl- $\Delta \gamma$ -dihydromuconate (XXV) To 0.74 g (0.0043 mole.) of the saturated lactonic ester (XXIV) in 8 ml of methanol was added 1.64 ml of methanolic sodium methoxide (2.62 N; 0.0043 mole.) at room temperature. After 15 min., the reaction mixture was evaporated under reduced pressure and then, were added 6 ml of water and it was acidified with dilute hydrochloric acid. An oily substance separated was extracted with ether. The solvent was removed and the residual product was completely

- 27 -

dried over paraffin in vacuo, n_D^{22} 1.4642, yield 1.20 g.

Ozonization: --- A 1.7 g quantity of the half-ester in chloroform was treated with an excess of ozone at 0° . The solvent was removed in vacuo, and the remaining ozonide was decomposed with water on a water bath for 10-15 min. and extracted with ether. The ethereal solution was distilled under reduced pressure to give methyl hydrogen oxalate (XXVII), bp. 110-4°/8 mm (Anschütz^{3,5)} bp. 108-9°/12 mm), yield 0.75 g. This half-ester was readily hydrolyzed to oxalic acid, mp. 186°, not depressed by admixture with an authentic specimen. The residue of the distillation was recrystallized from ethyl acetate, to yield methylsuccinic acid (XXVIII), mp. 106⁰, not depressed by mixed melting point comparison with an authentic specimen, 36) yield 0.9 g. This acid was also characterized by di-p-phenylphenacylester, mp. 179° (Anal. Found C 75.92, H 5.54, Calcd. for $C_{33}H_{28}O_6$ C 76.14, H 5.42).

 α -Methyl- $\Delta \gamma$ -dihydromuconic acid (XXVI) A l.17 g quantity of the half-ester (XXV) was warmed on a steam bath with 12.5 ml of l N sodium hydroxide for l hr., then, the reaction mixture was acidified with dilute hydrochloric acid, and extracted with ether, the acid was recrystallized from benzene in needles, mp. 121-2°. Anal. Found C 53.38, H 6.44, Calcd. for C₇H₁₀O₄ C 53.16, H 6.37. yield 0.64 g.

 (δ) -Methyl (a)-hydrogen a-methyl-cis,trans-muconate (XXIX) To the unsaturated lactonic ester (XXIII; 5.09 g, 0.0299 mole.) in 50 ml of methanol, was added 12.5 ml of methanolic sodium methoxide (2.39 N), After 15 min., the solution was evaporated under reduced pressure. Water (50 ml) was added and the solution was acidified with hydrochloric acid, whereupon methyl hydrogen a-methyl-cis,transmuconate separated, which crystallized from benzene in prismatic needles, mp. 121°; λ max. 275 m μ , ε 22,000. Anal.

- 28 -

Found C 56.53, H 5.98, Calcd. for $C_8H_{10}O_4$ C 56.46, H 5.92, yield 4.60 g.

Reconversion into the lactonic ester: — When the half-ester (0.27 g) was heated at $170-5^{\circ}$ under reduced pressure, a liquid distillate was obtained. After redistillation it gave the unsaturated lactonic ester (XXIII), bp. 150-5°/11 mm; n_D^{17} 1.4820 (yield 0.18 g), which was identified by the IR-spectrum.

a-Methyl-cis,trans-muconic acid (XVII c-t) A 1.50 g quantity of the preceding half-ester (XXIX) was warmed with 17.6 ml of l N sodium hydroxide on a steam bath for l hr. After the duration, the solution was acidified with hydrochloric acid, the separated crystal was recrystallized from methanol and from ethyl acetate to yield the cis,trans-acid in plates, mp. 172° , $\lambda \max$. 269 m μ , ϵ 25,700. Anal. Found C 53.87, H 5.23, Calcd. for $C_7H_8O_4$ C 53.84, H 5.16. yield 1.35 g.

Hydrogenation: — The hydrogenation of the cis,transacid (XVII c-t) over a platinum catalyst gave a-methyladipic acid (XXX), having mp. and mixed mp. 58-9°.

Methyl a-methyl-cis,trans-muconate (XXII c-t) Teatment of methyl hydrogen a-methyl-cis,trans-muconate (XXIX) with ethereal diazomethane, and evaporation of the solution afforded the dimethyl ester which, after recrystallization from methanol, formed prisms, mp. 60° ; λ max. 272 mµ, ε 27,600. Anal. Found C 58.84, H 6.70, Calcd. for $C_9H_{12}O_4$ C 58.69, H 6.57. The same dimethyl ester was also obtained by the same procedure from the cis,trans-acid (XVII c-t).

 α -Methyl-trans,trans-muconic acid (XVII t-t) (a) α -Methyl-cis,cis-muconic acid (XVII c-c; 0.51 g) was heated with 30 ml of 20% aqueous hydroxide for 4 hrs. The solution was cooled and acidified with dilute sulphuric acid. The separated precipitate was recrystallized from methanol to

- 29 -

yield the trans, trans-acid in prisms, mp. 273[°] (Kuhn and Michel²⁸⁾ recorded mp. 276[°]), $\lambda \max$. 273 mµ, ϵ 29,700. Anal. Found C 54.00, H 5.29, Calcd. for $C_7H_8O_4$ C 53.84, H 5.16. yield 0.45 g.

(b) α -Methyl-cis,trans-muconic acid (XVII c-t) gave the same acid exactly by the same procedures as mentioned above.

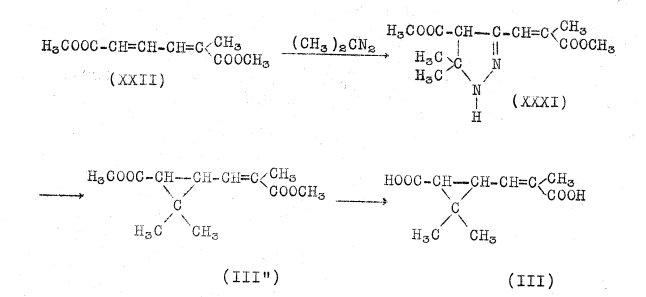
Hydrogenation of the trans, trans-acid gave α -methyladipic acid (XXX), as was the case with the cis, cis- and cis, trans-acids.

Methyl a-methyl-trans, trans-muconate (XXII t-t) The esterification of the trans, trans-acid (XVII t-t) with diazomethane gave the dimethyl ester in plates (from methanol), mp. 55.5°, $\lambda \max$. 275 mµ, ε 31,100 (Harris and Binns²⁹) reported mp. 46-7°, $\lambda \max$. 276 mµ, log ε 4.17), Anal. Found C 58.51, H 6.87, Calcd. for C₉H₁₂O₄ C 58.69, H 6.57.

V. ADDITION OF DIMETHYLDIAZOMETHANE TO METHYL α -METHYLMUCONATE¹¹)

Diemthyldiazomethane was added to methyl α -methyl-trans, trans-muconate (XXII t-t) in chilled xylene solution yielding an crystalline trans-pyrazoline ester (XXXI t), mp. 144-5°. Since the IR-spectrum of the pyrazoline ester indicated well-defined N-H (3270 cm⁻¹), carboxyl C=O (1712 cm⁻¹) and C=N (1675 cm⁻¹) bands as well as the UV-spectrum thereof had maxima at 213 m μ (ε 13,200; end absorption) and 299 m μ . (ε 9,600), the Δ^2 -pyrazoline structure was established for the pyrazoline ester. The addition of dimethyldiazomethane occurred, as was expected, selectively at the Yô-double bond of α -methylmuconate, this should reasonably be deduced from

- 30 -



the subsequent processes. The thermal decomposition of trans-pyrazoline ester (XXXI t) in the presence of copper as catalyst took place under vigorous evolution of nitrogen, and subsequent distillation in vacuo of the residue gave methyl trans, trans-chrysanthemumdicarboxylate (III" t-t), mp. $78-9^{\circ}$.

This crystalline ester as well as the corresponding free acid, mp. $206-8^{\circ}$, which was obtained by alkaline hydrolysis of the ester, were completely identical with the respective authentic specimens of methyl trans, trans-chrysanthemumdicarboxylate (III" t-t) and its free acid (III t-t) previously obtained through a different route by Inouye.² It was shown by the mixed melting point comparison as well

* As to the representation of configuration of chrysanthemumdicarboxylic acids and their esters, the first prefix indicates the geometry of the cyclopropane ring and then the second that of the side chain double bond, Thence, the order of those terms in the parent α -methylmuconic acids are reversed.

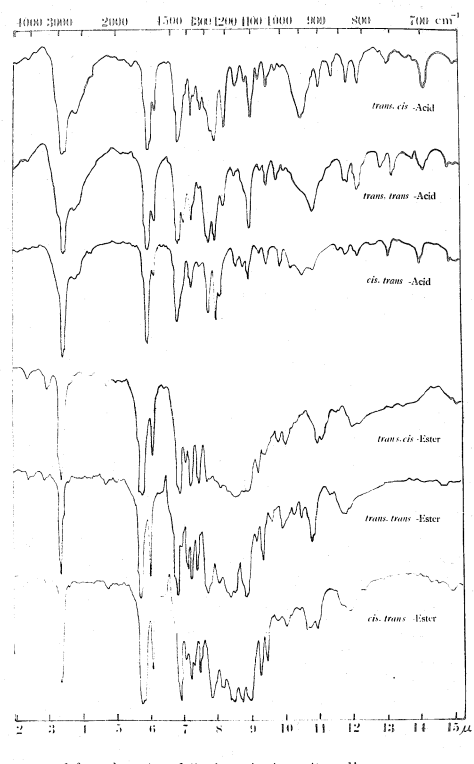
- 31 -

as by the complete identity of IR- and UV-spectra. The formation of trans, trans-chrysanthemumdicarboxylic acid (IIIt, t) from α -methyl-trans, trans-muconic acid (XVII t-t) by the addition of dimethyldiazomethane provides another evidence for the geometrical configuration of the side chain double bond of naturally occurring chrysanthemumdicarboxylic acid to be trans which has been concluded by the authors.^{8,9}

The addition of dimethyldiazomethane to methyl a-methylcis,trans-muconate (XXII c-t) gave cis-pyrazoline ester (XXXI c), mp. 99-100°. The $\gamma\delta - \varDelta^2$ -pyrazoline structure of this pyrazoline ester was also deduced from its IR-spectrum and the following procedures. The same cis-pyrazoline ester (XXXI c) was obtained by the addition of dimethyldiazomethane to methyl a-methyl-cis,cis-muconate (XXII c-c). It is reasonable that the same pyrazoline ester should be given from the cis,trans- and cis,cis-isomers of the starting material, because the configuration of $\alpha\beta$ -double bond of the starting materials are retained as such in the side chain double bond of the adducts and the specific geometry of the $\gamma\delta$ -double bond is broken, since the pyrazoline had a planar \varDelta^2 -pyrazoline structure.

On the thermal decomposition followed by hydrolysis, the cis-pyrazoline ester (XXXI c) converted into an acid of mp. 191° . By mixed melting point comparison with the authentic specimens of trans, trans- (III t-t) and cis, transchrysanthemumdicarboxylic acid (III c-t)²) respectively, this acid showed marked depression of the melting point. Then it was established that this acid differed from both trans, trans- and cis, trans-chrysanthemumdicarboxylic acids. Although the IR-spectra of this acid and its ester differed from those of trans, trans- and cis, trans-chrysanthemumdicarboxylic acids and their methyl esters in fine features, these were similar in most respects each other sufficient to

- 32 -



Infra-red spectra of the isomeric chrysanthemumdicarboxylic acids in nujol mull and their methyl esters in carbontetrachloride solution. establish that this acid had an analogous structure to these acids. The UV-absorption maximum of this acid (λ max. 236 m/) showed that this acid had the same chromophore as trans, trans- and cis,trans-chrysanthemumdicarboxylic acids, and its molecular extinction coefficient was consistent with the value expected by the cis-side chain structure. Therefore, this acid was established as trans,cis-chrysanthemumdicarboxylic acid (III t-c) having trans-structure about cyclopropane ring and cis form about side chain double bond.

Acid	mp.	λmax. ε	Dimethylester
trans, trans	206-8 ⁰	237 mji 15,400	mp. 78-9 ⁰
cis,trans	208-9 ⁰	234 14,700	bp. $100.5-101.5^{\circ}/$ $n_{\rm D}^{20}$ 1.4882
trans,cis	191 ⁰	236 9,500	bp. 101-2 [°] /1 mm n _D ²⁰ 1.4748

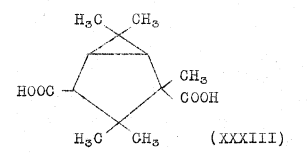
Matsui⁷⁾ have obtained an acid of mp. $175-7^{\circ}$ as trans, cis-chrysanthemumdicarboxylic acid starting from trans-caronylpropionitrile, however there yet remains some doubts in their conclusion that the trans,cis-isomer alone can be obtained stereospecifically in their processes, and also the mixture of equal amount of pure trans,trans- (mp. $206-8^{\circ}$) and trans,cis-acids (mp. 191°) prepared by the present author showed a similar melting point ($176-180^{\circ}$) to that of Matsui's acid. Therefore, the acid prepared by Matsui seems likely to contaminate the trans,trans-isomer. This was also convinced itself by Matsui and his coworkers, who described that they recognized their acid to be heterogeneous on X-ray diffractions and they tried to purify by means of

33

recrystallization but in vain.

It was somewhat surprising that trans, cis-chrysanthemumdicarboxylic acid retained its cis-configuration of the side chain double bond after being boiled in caustic alkaline solution, contrary to that the starting material, α methyl-cis, trans-muconic acid (XVII c-t), was easily isomerized under the same conditions. It seems likely to be related to the fact that chrysanthemumdicarboxylic acid is very difficult to add bromine as well as hydrogen.

We now return to the by-products in these processes. In the addition of dimethyldiazomethane to methyl α -methylcis.cis-muconate, a small amount of a pyrazoline ester of mp. 151⁰ (XXXII) was obtained, the yield of which varied from 5.5 to 0% according to the portion of dimethyldiazomethane employed. Both the analytical and spectral data showed that this pyrazoline ester should have a bicyclic pyrazoline structure which resulted from 1,4-addition of one molecule of dimethyldiazomethane and subsequent addition of another molecule of dimethyldiazomethane to the double bond newly created by the rearrangement. This pyrazoline ester gave an acid, mp. 220°, by thermal decomposition and subsequent hydrolysis, which is assumed to be 2,3,3,6,6-pentamethyl-bicyclo[3,1,0]hexane-2,4-dicarboxylic acid (XXXIII) from the analytical data but further elucidation of the structure has not yet been completed.



- 34 -

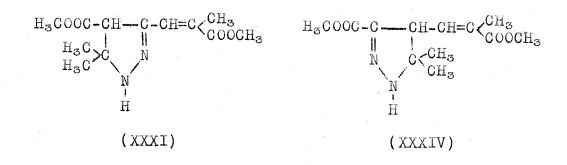
In the thermal decomposition of the pyrazoline esters, aliphatic by-products were obtained, and it was partially responsible for reducing the yield of the expected cyclopropane acids, chrysanthemumdicarboxylic acids.

From the trans-pyrazoline ester (XXXI t), a small quantity of an acid having mp. 121° was obtained as a by-product in the thermal decomposition and subsequent hydrolysis. This acid had a molecular formula $C_{10}H_{14}O_4$ identical with that of chrysanthemumdicarboxylic acid, and it showed Amax. at 218 m μ . The IR-spectrum of this acid resembled in most respects to that of the acid by-produced from cis-pyrazoline ester, described below, though there were some differences between them.

An acid of mp. 120-1° was obtained as a by-product in the thermal decomposition and subsequent hydrolysis of the cis-pyrazoline ester (XXXI c) in the yield of 16%. This acid had the same empirical formula $C_{10}H_{14}O_4$, as the trans-isomer mentioned above. The equivalent weight by titration of this acid was found as 191.8, which indicated that this acid had a formula $C_9H_{13}O_2COOH$; it was also supported by the analytical data of the amide derived therefrom. This acid absorbed 1 mole. equivalent of hydrogen over a platinum catalyst giving an acid of mp. $\frac{13.5}{130-1}$, which had an empirical formula $C_{10}H_{16}O_4$. Therefore, it was expected that the acid of mp. 121° was an acyclic isomeric acid which formed on the way of thermal decomposition of the pyrazoline ring, and one of the two carboxylic groups became bound with one of the double bonds as to form a lactone ring. This expectation was supported by the fact that this acid consumed 2 mole of alkali after being refluxed with alkaline solution and that the IR-spectra of this acid and of its amide showed an lactonic C=0 band at 1718 cm^{-1} and 1678 cm^{-1} respectively. These lactonic C=O bands were located in longer wave length

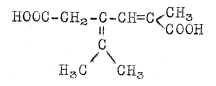
- 35 -

reagion than those of Y- or δ -lactones except $\alpha\beta$ -unsaturated δ -lactone,³⁷⁾ then this acid seems to have the $\alpha\beta$ -unsaturated δ -lactone or less strained large lactone ring. The UV-spectrum of this acid showed an end absorption maximum at 220 mµ and did not show any other maximum, that indicated that this acid had not more-conjugated unsaturated systems than $\alpha\beta$ -unsaturated carboxyl system. Regarding the structure of the parent substance (XXII), two structures are considerable to the pyrazoline ester in β^2 -form, i.e. in one of them the terminal nitrogen atom of dimethyldiazomethane is bonded

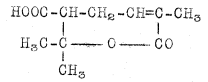


with the Y-carbon atom of α -methylmuconate, as shown in formula XXXI, and in the other the nitrogen becomes bound at δ carbon atom of α -methylmuconate, as shown in formula XXXIV. However, the latter form is excluded by the mechanistic argument of the addition of diazocompound mentioned in chapter III, and also by the fact that the acyclic isomer (XXXV) expected from the pyrazoline structure (XXXIV) was already obtained by Inouye² which was too stable to lactonize under the conditions used. Therefore, the structure of this byproduced acid might be α & -trimethyl- δ -carboxy- $\Delta \alpha$ -hexenolide (XXXVI).

- 36 -



(XXXV)



(XXXVI)

EXPERIMENTAL

Addition of dimethyldiazomethane to methyl $\alpha\text{-methyl-muconates.}$

Addition to the trans, trans-ester. (a) To the dimethyldiazomethane solution in xylene prepared from 5.3 g of acetone-hydrazone by Guha and Sankaran's method, were added 4.9 g of methyl a-methyl-trans, trans-muconate (XXIIt-t) under vigorous stirring and cooling at $-15 - -20^{\circ}$. The cooling and stirring were continued for ca. 3 hrs., and then the solution was stood still overnight at room temperature. The reaction mixture was separated from a slaggy mercury and mercurous oxide by filtration. The filtrate was concentrated under reduced pressure. By cooling, the crystal of transpyrazoline ester (XXXI t) was obtained in lathes (from methanol or ethyl acetate), mp. 144-5° (decomp.), yield 3.7 g (56%). Anal. Found C 56.59, H 7.01, Calcd. for C₁₂H₁₈O₄N₂ С 56.68, Н 7.14.

IR-spectrum: N-H 3270, carboxyl C=0 1712, C=N 1675 cm⁻¹. UV-spectrum: λmax. 213 mμ (end absorption), ε 13,200; 299 mμ, ε 9,600

(b) Addition to the cis, trans-ester. To the dimethyldiazomethane solution prepared from 37 g acetone-hydrazone,

- 37 -

were added 31.4 g of methyl α -methyl-cis,trans-muconate (XXII c-t) in the same condition as the trans,trans-isomer. Xylene was removed from the reaction mixture and the crystal obtained by cooling of the residue was recrystallized from methanol yielding the cis-pyrazoline ester (XXXI c), mp. 99-100[°] (decomp.), plates, yield 21.4 g (49%). Anal. Found C 56.65, H 7.27, Calcd. for $C_{12}H_{18}O_4N_2$ C 56.68, H 7.14.

IR-spectrum: N-H 3230, carboxyl C=0 1715, C=N 1672 cm⁻¹ UV-spectrum: λmax. 213 mμ (end absorption), ε 10,700; 299 mμ, ε 10,200

Addition to the cis, cis-ester. The addition of (c) dimethyldiazomethane, prepared from 35.0 g (3 mole. equivalent to the olefinic compound), to 30.0 g of methyl α methyl-cis, cis-muconate (XXII c-c) was pursued in the same way as above-mentioned. As well as the cis-pyrazoline ester (XXXI c), the other pyrazoline ester of mp. 151° (decomp.) (XXXII) was obtained in lathes. The yield of them were 17 g (41%) and 2.3 g (5.6%) respectively. In the case that the ratio of acetone-hydrazone used for preparing dimethyldiazomethane and methyl a-methyl-cis, cis-muconate was 1.5 to 1.0, the by-product of mp. 151° could not be obtained, though there is no distinct variation in the yield of the cispyrazoline ester (XXXI c). For the structure of the latter adduct, see the text. Anal. Found C 55.65, H 7.74, N 17.09, Calcd. for $C_{15}H_{24}O_4N_4$ C 55.54, H 7.46, N 17.27. UV-spectrum: λ max. 214 mµ (end absorption), ϵ 3,580.

Thermal decomposition of the pyrazoline esters.

(a) Decomposition of the trans-pyrazoline ester. One gram of the trans-pyrazoline ester (XXXI t) was mixed with 0.1 g of Gattermann's copper powder and heated at 160-170°, whereby the decomposition occurred with evolution of

- 38 -

nitrogen. After nitrogen release was over, copper and resinous product insoluble in petroleum ether were removed, and the resulting oily substance was distilled under reduced pressure. Bp. $104-8^{\circ}/1 \text{ mm}$; $n_{\tilde{D}}^{20}$ 1.4855; yield 0.71 g (79%). The distillate was partly crystallized after repeated rectifications, mp. 78-9°. The melting point was not depressed by the mixed melting point comparison with an authentic specimen of methyl trans, trans-chrysanthemumdicarboxylate (III" t-t)?

(b) Decomposition of cis-pyrazoline ester. Five grams of cis-pyrazoline ester (XXXI c) was decomposed with 0.5 g of copper powder at $160-170^{\circ}$. The decomposition product was treated as above-mentioned, and an oily substance was obtained. Bp. $103-5^{\circ}/1$ mm; $n_{\rm D}^{20}$ 1.4758; yield 3.1 g (70%).

Hydrolysis of the esters (III")

A 0.58 g quantity of the ester of bp. $104-8^{\circ}/1$ mm (a) was refluxed with 5.5 ml of 5% methanolic sodium hydroxide on a steam bath for 2 hrs. After removal of the solvent, the residue was dissolved in water, acidified with hydrochloric acid and extracted with ether. The ethereal extract was recrystallized from ethyl acetate to give the expected trans.trans-chrysanthemumdicarboxylic acid (III t-t), mp. 206-8° (mixed melting point with an authentic specimen²) was 206-8°), yield 0.35 g (69%), as well as an by-produced acid of mp. 122°, yield 0.07 g (14%). The characteristics of the latter acid, Anal. Found C 60.90, H 7.40, Calcd. for C 60.59, H 7.12. UV-spectrum: $\lambda \max$. 218 mµ, ϵ $C_{10}H_{14}O_{4}$ 14,200.

(b) In exactly the same way as mentioned above, 3.0 g of the ester of bp. $103-5^{\circ}/1$ mm was hydrolyzed. trans,cis-Chrysanthemumdicarboxylic acid (III t-c), mp. 191° , and an

- 39 -

acid of mp. $120-1^{\circ}$, were obtained after fractional recrystallization from chloroform or methanol. The yield of them were 0.6 g (23%) and 0.4 g (16%) respectively. The characteristics of trans, cis-chrysanthemumdicarboxylic acid were as follows, prisms (from methanol), mixed melting points with trans, trans-chrysanthemumdicarboxylic acid (mp. 206-8°) and the cis, trans-isomer (mp. 208-9°) were 176-80° and 167-72° respectively. Anal. Found C 60.68, H 7.38, Calcd. for $C_{10}H_{14}O_4$ C 60.59, H 7.12.

IR-spectrum: in the text.

UV-spectrum: $\lambda \max$. 236 m μ , ϵ 9,500

This acid was recovered unchanged after being refluxed with aquious 20% sodium hydroxide for 4 hrs., mp. 191° (recrystallized from ethyl acetate).

This acid was esterified with diazomethane to yield quantitatively its methylester (III" t-c), bp. $101-2^{\circ}/1$ mm; n_D^{20} 1.4748.

The data of the byproduced acid of mp. $120-1^{\circ}$ (XXXVI) were mentioned below. Prisms (from methanol). Anal. Found C 60.78, H 7.24, Calcd. for $C_{10}H_{14}O_4$ C 60.59, H 7.12. Eq.wt. Found 191.8, Calcd. for $C_{9}H_{13}O_2$ COOH 198.21. UV-spectrum: λ max. 220 mµ (end absorption), ε 6,700

Hydrogenation: — A 0.998 g quantity of the acid of mp. $120-1^{\circ}$ (XXXVI) in 80 ml of methanol was hydrogenated over a platinum catalyst (PtO₂ 30 mg) in a shaking apparatus and absorbed 125 ml (at 19°) of hydrogen, equivalent to one double bond in $C_{10}H_{14}O_4$. The reduction product was freed from catalyst and the solvent, and recrystallized from benzene to give an acid , mp. 132.5-133.5°, long needles, Anal. Found C 60.10, H 8.15, Calcd. for $C_{10}H_{16}O_4$ C 59.98, H 8.05. Eq.wt. Found 201.2, Calcd. for $C_{9}H_{15}O_2COOH$ 200.23, eq.wt. after the acid was refluxed with 1 N sodium hydroxide on a steam bath for 1 hr., Found 104.2, Calcd. for

- 40 -

 $C_8H_{16}O(COOH)_2$ 109.12. Amide, mp. 152.5-3^o, needles (from chloroform and light petroleum), Anal. Found C 60.91, H 7.66, N 7.40, Calcd. for $C_{10}H_{15}O_3N$ C 60.89, H 7.67, N 7.10.

Thermal decomposition and subsequent hydrolysis of the pyrazoline ester (XXXII). A 2.41 g quantity of the pyrazoline ester (XXXII), mp. 151°, was decomposed with 0.15 g of copper powder, to give an oily substance, bp. $111-3^{\circ}/0.5$ mm, n_D^{20} 1.4688, yield 1.48 g. This ester (1.48 g) was hydrolysed and the product was recrystallized from ethyl acetate or benzene yielding an acid (XXXIII) of mp. 218-20°. Anal. Found C 65.66, H 8.51, Calcd. for $C_{1.3}H_{20}O_A$ C 64.98, H 8.39.

REFERENCES

- H.Staudinger, L.Ruzicka: Helv.Chim.Acta, 7,177 (1924)
 a) Y.Inouye, Y.Takeshita, M.Ohno: Bull.Agr.Chem.Soc.
 - Japan, <u>19</u>,193 (1955); Bull.Inst.Chem.Res.Kyoto Univ., <u>33</u>,73 (1955)

b) Y.Inouye, M.Ohno: Bull.Agr.Chem.Soc.Japan, <u>21</u>,265 (1957); Bull.Inst.Chem.Res.Kyoto Univ., <u>34</u>,90 (1956)

- (3) S.H.Harper, K.C.Sleep: Chem.Ind., <u>1954</u>,1538;
 L.Crombie, S.H.Harper, K.C.Sleep: J.Chem.Soc., <u>1957</u>, 2743
- (4) Y.Fujitani: Arch.Exper.Path.Pharm., <u>61</u>,47 (1909)
- (5) L.Crombie, S.H.Harper: J.Chem.Soc., <u>1954</u>,470
- (6) Y.Inouye, M.Ohno: Kagaku (Tokyo), <u>28</u>,636 (1958)
- M.Matsui, M.Miyano, Y.Yamashita, H.Kubo, K.Tomita,
 Bull.Agr.Chem.Soc.Japan, <u>21</u>,22 (1957)
- (8) Y.Inouye, T.Sugita, M.Ohno: Bull.Agr.Chem.Soc.Japan, <u>21</u>,5 (1957)

- 41 -

(9)	Y.Inouye, T.Sugita, M.Ohno: Bull.Agr.Chem.Soc.Japan,
	<u>21</u> ,222 (1957)
(10)	Y.Katsuda, T.Chikamoto, Y.Inouye: Bull.Agr.Chem.Soc.
	Japan, <u>23</u> ,174 (1959)
(11)	S.Takei, T.Sugita, Y.Inouye: Ann., <u>618</u> ,105 (1958);
	Y.Inouye, T.Sugita, M.Ohno: Bull.Agr.Chem.Soc.Japan,
	<u>22</u> ,269 (1958)
(12)	A.v.Baeyer, H.Rupe: Ann., <u>256</u> ,1 (1890)

- (13)H.Burton, C.K.Ingold: J.Chem.Soc., 1929,2022
- (14)C.K.Ingold; Structure and Mechanism in Org. Chem., 696
- E.H.Farmer, L.A.Hughes: J.Chem.Soc., 1934,1929 (15)
- (16)Barbier, R.Locquin: Compt.rend., 156,1445 (1913)
- P.Heinänen: Suomen Kemistilehti, 11, B,2 (1938); Chem. (17)Zent., 1938,4032
- Ann., <u>353</u>,178 (1907) (18)R.Anschütz:
- (19)Org.Synth., Coll.Vol. II, 382

- (20)T.Sugita, Y.Inouye, M.Ohno: Bull.Agr.Chem.Soc.Japan, 22, 162 (1958)
- (21)N.L.Drake, J.Bronitsky: J.Am.Chem.Soc., <u>52</u>,3715 (1930)
- (22)Y.Inouye, M.Ohno: Botyu-Kagaku, 20,136 (1955)
- (23)Y.Inouye, M.Ohno: Bull.Agr.Chem.Soc.Japan, 20,77 (1956)
- (24)S.H.Harper, H.W.B.Reed: J.Chem.Soc., 1955,779
- (25)H.M.Walborsky: private letter.
- C.v.Heide: Ber., 37,2101 (1904) (26)
- P.C.Guha, D.K.Sankaran: Ber., 70,1688 (1937) (27)
- (28)R.Kuhn, J.Michel: Ber., 71,1119 (1938)
- J.O.Harris, F.Binns: Nature, 179,475 (1957) (29)
- (30)J.A.Elvidge, R.P.Linstead, P.Sims, B.A.Orkin: J. Chem. Soc., 1950,2235
- (31)J.A.Elvidge, R.P.Linstead, J.F.Smith: J.Chem.Soc., 1952,1026

- 42 -

- (32) J.A.Elvidge, R.P.Linstead, P.Sims: J.Chem.Soc., <u>1951</u>, 3386
- (33) F.P.Mazza, G.DiMase: Gazz.chim.ital., <u>57</u>,300 (1927)
- (34) L.Bouveault, R.Locquin: Compt.rend., <u>146</u>,138 (1908); Bull.Soc.Chem.France, [4]<u>3</u>,451 (1908)
- (35) R.Anschütz: Ann., 254,1 (1889)
- (36) Org.Synth., Coll.Vol. III, 615
- (37) F.Korte, K.H.Büchel, K.L.Göhring: Angew.Chem., <u>71</u>,523 (1959)
- (38) P.C.Guha, D.K.Sankaran: Ber., <u>70</u>,1688 (1937)

ACKNOWLEDGEMENT

The author wishes to express his sincere appreciation to Professor M.Ohno and Dr. Y.Inouye for their encouragement and guidance throughout the course of this investigation.

He also expresses his deep gratitude to Professor S.Takei and Professor M.Nakajima for many helpful suggestions.