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The Chemistry of 1,5,9-Cyclododecatriene and Syntheses Based on It

L.I.Zakharkin and V.V.Guseva

The methods of synthesis and chemical properties of 1,5,9-cyclododecatriene (the butadiene cyclotrimerisation product), which is manufactured on an industrial scale, are examined; syntheses based on this interesting compound have played a major role in the development of organic chemistry of medium and large rings. The bibliography includes 406 references.

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I. INTRODUCTION

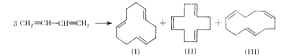
The first studies on cyclododecane and its derivatives were begun in 1926 by Ruzicka and coworkers, but the difficulty of the synthesis of these compounds and of their isolation in a pure form hindered the development of their chemistry.

In 1954, Reed reported for the first time¹ the synthesis of a cyclic butadiene trimer in the presence of bis(triphenylphosphine)nickel dicarbonyl, but it was not until Wilke's study² of the cyclotrimerisation of butadiene in the presence of organometallic complex catalysts that 1,5,9-cyclododecatriene (1,5,9-CDT) became readily available and vigorous development of the chemistry of this compound and its industrial use were initiated.

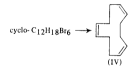
cis, trans, trans-1,5,9-CDT is manufactured at present on a large scale in many countries. The following products obtained from 1,5,9-CDT are of major technical importance: ω -dodecalactam—the monomer for the synthesis of nylon-12; decane-1,10-dicarboxylic acid—the monomer for the synthesis of polyamides and polyesters and a substitute of sebacic acid in the synthesis of plasticisers and lubricating oils; 1,12-diaminododecane—the monomer for the synthesis of polyamides; 1,2,5,6,9,10hexabromocyclododecane, which is used as an additive to polymers in order to render them incombustible; cyclododecanone which is used as the initial compound for the synthesis of valuable perfumes—cyclopentadecanone (exaltone) and the lactone of ω -hydroxypentadecanoic acid (tibetolide).

II. SYNTHESIS AND CHEMICAL REACTIONS OF 1,5,9-CYCLODODE CATRIENE ISOMERS

It has now been established that the cyclotrimerisation of butadiene to 1,5,9-CDT proceeds in catalytic systems containing the following transition metals as components: titanium, chromium, nickel, and manganese. *trans*, *trans*, *trans*-1,5,9-CDT (I), *cis*, *trans*, *trans*-1,5,9-CDT (II), and *cis*, *cis*, *trans*-1,5,9-CDT (III) have been obtained in the presence of these catalysts; the ratios of the yields of these products depend on the nature of the transition metal, the ligands in the catalytic system, and the reaction conditions:



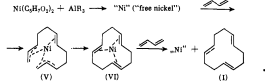
The fourth possible isomer—cis, cis, cis, 1,5,9-CDT (IV)—is not formed in the catalytic trimerisation of butadiene. It has been obtained by the dehydrobromination of hexabromocyclododecane—the product of the addition of three bromine molecules to (I):³



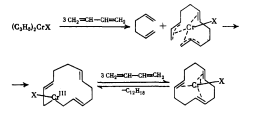
The most effective catalysts for the synthesis of 1,5,9-CDT are systems containing various titanium compounds $[TiCl_4, {}^{4-20} TiBr_4, {}^{21} Ti(OR)_4, {}^{22,23} (RO)_2 TiO, {}^{24} TiCl_{4-n}(OR)_n, TiCl_{4-n}(OCOR)_n, {}^{55-28}$ and the complex C_6H_6 .TiCl_2.Al_2Cl₆ 13 , 29,30] in combination with reducing agents such as trialkylaluminium or aluminium alkyl halides, calcium hydride, and dialkylaluminium hydride. The reaction proceeds only in solution in aromatic hydrocarbons with formation of 1,5,9-CDT in 80-90\% yield. The 1,5,9-CDT obtained is a mixture of two isomers: 3-4% of (I) and 96-97% of (II). A mixture containing 67% of (I) and 33% of (II) is formed in the catalytic system containing poly(butyl orthotitanate) and $(C_2H_5)_2AlCl.^{31}$

Catalytic systems containing chromium and manganese $[(C_5H_7O_2)_2Cr, CrO_2Cl_2 \text{ or } CrO_3 + (C_2H_5)_3Al,^{4,5,12} CrCl_3 + (iso-C_4H_9)_3Al,^{10-12,32,33} and (C_5H_7O_2)_2Mn + (C_2H_5)_3Al^{34}]$ lead to a mixture of approximately 60% of (I) and approximately 40% of (II). A mixture consisting of 65–80% of (I), 7–9% of (II), 9–11% of (III), 7–9% of vinylcyclohexene, and 4–6% of 1,5-cyclo-octadiene is formed in the presence of nickel catalysts [Ni(C_5H_7O_2)_2 + C_2H_5OAl(C_2H_5)_2, 12,35,36] NiX_2 + (iso-C_4H_9)_3Al,^{37} (cyclo-octadiene)_2.Ni(O) ^{30,36} and cyclodode catriene.Ni(O) ³⁵]. The mixture composition varies with reaction temperature.

There have been numerous patents for catalytic systems containing nickel^{37,38-43}. The mechanism of the reaction in the presence of nickel catalysts has been established by Wilke and coworkers^{33,35,44-46}. It includes the following stages:



When Ni($C_5H_7O_2$)₂ is reduced by an organoaluminium compound, "free" nickel is formed, which trimerises butadiene to the intermediate π -allyl complex of nickel (V); ring closure of (V) results in the formation of the π -nickel complex (VI). Treatment with butadiene leads to the displacement of 1,5,9-CDT from this complex. Complexes (V) and (VI) have been isolated and their structures have been determined by X-ray diffraction^{47,48} and confirmed by certain reactions^{31,55,44,45}. When complex (VI) is acted upon by cis, cis, cis-1,5,9-CDT (IV), trans, trans, trans-1,5,9-cyclododecatriene (I) is displaced from it and the complex (IV).Ni(O) is formed⁴⁵; the latter readily reacts with carbon monoxide to give Ni(CO)₄ and compound (IV). A mechanism has been proposed for the trimerisation of butadiene in the presence of chromium catalysts³³ in which the valence state of chromium changes continuously from +1 to +3:



It has been shown 30,49 that in the presence of titanium catalysts the active centre for the trimerisation of butadiene is a complex in which titanium is present in the bivalent state:

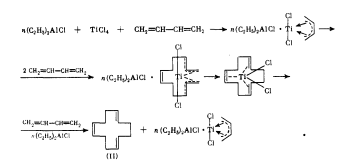


Table 1 presents the best known catalytic systems in the presence of which the trimerisation of butadiene to 1,5,9-CDT isomers takes place with a high yield.

Among the four 1,5,9-CDT isomers, the trans, trans, trans-isomer (I)⁵⁰ and the cis, trans, trans-isomer (II), which are readily obtainable in the presence of the above catalysts, have been studied in greatest detail. The presence of three non-conjugated double bonds in the 1,5,9-CDT molecule makes this system capable of undergoing many reactions. Among them the most interesting are those involving cis-trans isomerisation. When (II) is illuminated with ultraviolet light, a mixture of (I) and (III) is formed ⁵¹.

Table 1.

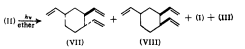
Catalytic systems	Yield of 1,5,9-CDT, %	Isomers formed (the relative percentage contents are indi- cated in brackets)	Refs.
$\begin{array}{l} TiCl_4 + (C_8H_8)_2AlCl \\ TiCl_4 + (C_8H_8)_2Al_3Cl_3 \\ CrO_9Cl_3 + (C_8H_9)_2Al_3Cl_3 \\ CrCl_9 + (C_8H_9)_2Al_3 \\ CrCl_9 + (C_9H_8)_2Al_3 \\ CrCl_9 + (so-C_4H_8)_2Al_3 \\ Ni(C_4H_7O_8)_2 + (C_8H_8)_2Al_3 \\ NiX_8 + (so-C_4H_8)_2Al_3 \\ [(C_4H_{12})_8Nl] \\ \end{array}$	91 88 70 80 86 81 80 87	$ \begin{vmatrix} f(97) + II & (3) \\ II & (98) + I & (2) \\ I & (59) + II & (41) \\ I & + II \\ I & (60) + II & (40) \\ I & (80) + II & (9) + III & (11) \\ I & + II \\ I & + II \end{vmatrix} $	8, 9, 10 8, 9, 12 10 8 8, 9 10 37 39

Subsequently it was shown that each of the four isomers undergoes cis-trans isomerisation giving rise to the same equilibrium mixture consisting of all four isomers (I), (II), (III), and (IV).⁵² It has been established⁵³ that thermo-dynamic stability decreases in the sequence (I) > (II) > (III). Data for the photochemical isomerisation of the isomers in benzene are presented in Table 2.

Table 2.Composition of the mixture of 1,5,9-CDT isomers in photochemical isomerisation

Initial	Irradiation time, h	Proportions of isomers in product, %		
olefin		(I)	(11)	(111)
(I)	17	24	43 55	33
(II) (III)	17 100	11	55 21	33 34 78

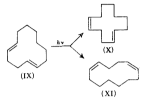
Table 2 shows that (III) isomerises much more slowly than isomers (I) and (II). The proportions of the isomers also depend on the selected sensitiser. In the presence of aromatic ketones and quinones, isomer (II) predominates in the resulting mixture, while the use of aliphatic ketones leads to isomer (III). Thus the readily available isomer (II) may be easily converted into (I) and (III) on irradiation of a benzene solution containing acetophenone or acetone. After prolonged irradiation of a mixture of (I), (II), and (III), 43% of trans, trans-2,6-divinyl-cis-bicyclo[3,3,0]octane was obtained together with 1,5,9-CDT isomers⁵². When (II) in an ethereal solution is irradiated, a mixture containing 25% of (I), 29% of (III), and isomeric 1,2,4-trivinylcyclohexanes [12% of (VII) and 1% of (VIII)] is formed⁵⁴:



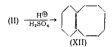
Thermal isomerisation of (II) at 450° C leads only to isomeric 1,2,4-trivinylcyclohexanes⁵⁵. In the presence of toluene-*p*-sulphinic acid a mixture of 80-85% of (I) and 15-20% of (II) is formed from the initial (I)-(III) mixture on

heating to 90° C.^{53,56} Copper salts are active catalysts of the *cis-trans* isomerisation of the double bonds in (I), (II), and (III).⁵³ The free-radical isomerisation of (II) to (I) takes place readily in the presence of thioglycolic acid⁵⁷, benzoyl peroxide⁵⁷, and *N*-bromosuccinimide^{57,58}. The photochemical isomerisation of cyclododecenes in benzene leads to the formation of the less stable *cis*-isomer⁵³. In the presence of copper salts the equilibrium is also displaced to *cis*-cyclododecene⁵³.

On photochemical isomerisation in benzene, *trans*, *trans*-1,5-cyclododecadiene (IX) is converted into the *cis*, *trans*-isomer (X) and the *cis*, *cis*-isomer (XI); the ratios of the isomers in the resulting mixture are (IX):(X):(XI) =17:55:28⁵⁹:

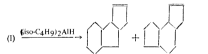


In the presence of acid reagents (II) undergoes intramolecular cyclisation, which leads to products with bicyclic and tricyclic structures. Thus a mixture of hydrocarbons in which $\Delta^{1,6}$ -bicyclo[4,6,0]dodecene (XII) predominates, is formed from (II) on treatment with 80% sulphuric acid in the cold⁶⁰:

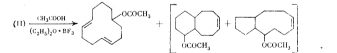


A mixture of acenaphthene and decahydroacenaphthene is formed on heating (II) with polyphosphoric $acid^{81}$.

The isomerisation of (I) takes place readily in the presence of $(iso-C_4H_9)_2$. AlH on heating to 200°C, giving an 85% yield of a mixture of tricyclo[6,4,0,0^{3,7}]dodec-4-ene and tricyclo[7,3,0,0^{3,7}]dodec-4-ene⁶²:



Bicyclic products were formed after the addition of acetic acid and hydrogen chloride to $(\Pi)^{\,63}$:

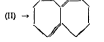


The reaction apparently proceeds both at the cis-double bond, giving rise to the usual addition products, and at the *trans*-double bond, giving rise to bicyclic products.

When a cetic acid and hydrogen chloride react with (I), only addition products retaining the molecular skeleton are formed 63 :



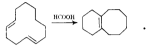
Bicyclo[5,5,0]dodeca-1,7-diene is formed from (II) in the presence of Na/Al₂O₃, 64 Co₂(CO)₈, 65 and the lithium derivative of ethylenediamine 66 :



When methyl trans, trans-4,8-cyclododecadiene-carboxylate is treated with 90% formic acid, 2,3,6,7,8,9-hexahydro-1H-benz[e]indene is produced⁶⁷.

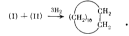


trans, trans-1,5-cyclododecadiene gives rise to bicyclo-[6,4,0]dodecene⁶⁷:



III. SYNTHESES BASED ON cis, trans, trans- AND trans, trans, trans-1,5,9-CYCLODODE CATRIENES

In this section of the review we shall consider the reactions of the most readily available 1,5,9-CDT isomers the *trans*, *trans*, *trans*-isomer (I) and the *cis*, *trans*, *trans*isomer (II). The majority of studies on the hydrogenation of 1,5,9-CDT in cyclododecane are described in the patent literature. The two isomers (I) and (II) are quantitatively hydrogenated in the presence of platinum, palladium, and Raney nickel⁶⁸⁻⁷⁰:



In order to obtain large amounts of cyclododecane, (I) and (II) are hydrogenated at elevated temperatures in the presence of Ni/Al₂O₃ or Ni/Cr₂O₃. ^{5,68-72} Nickel, cobalt, and copper salts deposited on SiO₂ or Al₂O₃ in the presence of NaF are effective catalysts of the reduction ^{73,74}. The rates of hydrogenation of the two isomers are virtually the same. In the presence of these catalysts, all three double bonds are hydrogenated at virtually the same rates and one cannot therefore carry out a selective hydrogenation. When one mole of (II) is hydrogenated with two moles of hydrogen in the presence of palladium, a mixture consisting of cyclododecene, cyclod decadiene, and cyclododecanes is formed ^{68,75}. The hydrogenation of (I) and (II) to cyclododecane in the presence of homogeneous catalysts has been described in a number of patents: $(Ph_3As)_2(CO)RhCl + AlCl_3, ⁷⁶ Ni(C_5H_7O_2)_2 + LiAlH_2(OCHMeC_2H_5)_2, ⁷⁷ Ti(OBu)_4 + (C_2H_5)_2AlCl, and Cr(C_5H_7O_2)_2 + (C_2H_5)_2AlCl + (C_2H_5)_3Al.⁷⁹$

The selective hydrogenation of (I) and (II) to cyclododecene, which gives rise to great possibilities for the synthesis of many products of practical importance, is of considerable interest. It has been shown that, when (II) is acted upon by di-imide, the *trans*-double bonds are reduced initially and *cis*-cyclododecene is formed in a high yield 80,81 :

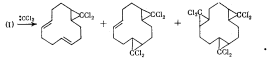


The success of the catalytic hydrogenation of (II) to cyclodode cene depends primarily on the nature of the catalyst. The use of relatively inactive heterogeneous catalysts makes it possible to stop the hydrogenation at the stage corresponding to the addition of two moles of hydrogen⁸²⁻⁸⁷. Studies on a series of catalysts (Rh, Pd, and Ru on Al₂O₃)^{88,89} have shown that the hydrogenation of (I) and

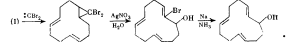
(II) to cyclododecene takes place most selectively in the presence of rhodium. The catalysts can be arranged in the following sequence in terms of decreasing yields of cyclododecene: Ph > Pd > Ru > Pt. The trichlorotripyridinerhodium +NaBH₄ catalytic system proved to be the most effective in the hydrogenation (II) to cyclododecene⁹⁰. In the presence of the homogeneous catalyst [Co(CO)₃. PR₃]₂ ^{83,964,91} or (R₃P)₂(CO)₂RuCl₂ ^{92,93} (II) gives a high yield of a mixture of *cis*- and *trans*-cyclododecenes (96–98%) and in addition 1% of 1,5-cyclododecadiene and 0.5% of cyclododecane are formed. Hydrogenation of (II) in the presence of 7% PtCl₂ in a (C₂H₅)₄NSnCl₂ melt at 160°C and 100 atm of H₂ leads to 2% of (II), 10% of cyclododecadiene, and 87% of cyclododecene⁹⁴.

The addition of halogens and halogenocarbenes, oxidation, hydroboration, and carboxylation are other reactions involving double bonds in (I) and (II). When (I) and (II) are brominated, a mixture of dibromocyclododecadiene, tetrabromocyclododecene, and hexabromocyclododecane is formed 68,95,96 . However, it has been stated in patents 97,98 that, when one mole of (I) is brominated with two moles of Br₂ in the presence of sodium acetate, tetrabromocyclododecene is formed quantitatively. It has been shown⁶⁸ that hexabrom ocyclodode cane obtained from (II)has a melting point of $194-195^{\circ}C$, while that obtained from (I) has a melting point of $177-178^{\circ}C$. These brom o-derivatives are different geometrical isomers. On recrystallisation from benzene, the hexabromocyclododecane obtained from (I) gives rise to the clathrate compound $(C_{12}H_{18}Br_6)_2C_6H_6$.⁷⁵ Hexabromocyclododecane has four Hexabromocyclododecane has found an application as an additive to polymeric materials in order to render them incombustible.

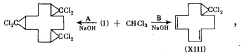
The addition of halogenocarbenes to (I) and (II) has been investigated in detail. The addition of : CCl_2 to (II) takes place preferentially at the *trans*-double bond⁹⁹. When :CHCl¹⁰⁰ and :CCl₂¹⁰¹ act on (I), a mixture of three products is formed:



Treatment of (I) with : CBr_2 gives a high yield of dibromobicyclotridecadiene¹⁰², which on treatment with AgNO₃ in water is converted into bromohydroxycyclotridecatriene; on treatment with sodium in liquid ammonia, the latter is converted into hydroxycyclotridecatriene:



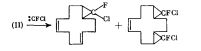
The addition of $:CCl_2$ to (II) under the conditions of phase-transfer catalysis leads to different results depending on the ammonium salt used ¹⁰³:



where $A = \text{cetyltrimethylammonium bromide and } B = \text{tri-alkyl-}\beta-\text{hydroxyethylammonium bromide}$. When the diene (XIII) is hydrogenated, the product is 13,13-dichlorobi-cyclo[10,1,0]tridecane, from which cyclotridecanone was obtained ¹⁰⁴:

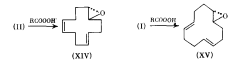
$$(XIII) \xrightarrow{H_2} (CH_2)_{10} \xrightarrow{CH} CCl_2 \xrightarrow{\mathbf{n} \cdot C_4 H_9 L_1} (CH_2)_{10} \xrightarrow{CH} (H_2SQ_4 (CH_2)_{11} \xrightarrow{C=0} (CH_2)_{11} \xrightarrow{C=0} (CH_2)_{11} \xrightarrow{C=0} (CH_2)_{11} \xrightarrow{C=0} (CH_2)_{11} \xrightarrow{CH} (CH_2)_{12} \xrightarrow{CH} (CH_2)_{12} \xrightarrow{CH} (CH_2)_{13} \xrightarrow{CH} (CH_2)_{14} \xrightarrow{CH} (CH_2)_{1$$

The reaction of (II) with :CFCl takes place at the *trans*double bonds with formation of two addition products 105 :



The addition of :CHOCH₃ to (I) entails the formation of only trans-13-methoxybicyclo[10,1,0]trideca-trans, trans-4,8diene, while a mixture of diastereoisomers is formed from (II) under these conditions ¹⁰⁵. The reaction of :CCl₂ with cis, trans-1,5-cyclododecadiene takes place unambiguously, resulting in the formation of trans-13,13-dichlorobicyclo-[10,1,0]tridec-4-ene(cis). ¹⁰⁸ A study of the rates of addition of :CHCl, ¹⁰⁰ :CBr₂, ¹⁰⁷ and :CHOCH₃ ¹⁰⁶ to cis- and trans-cyclododecenes showed that the reaction involving the trans-double bond is faster.

The oxidation of (I) and (II) to epoxycyclododecadiene, further transformations of which lead to interesting and valuable products, is preparatively important. The epoxidation of (I) and (II) has been carried out with peracetic, perbenzoic, and perphthalic acids, as well as a mixture of H_2O_2 and CH_3COOH in the presence of a cationexchange resin^{5,11,68,109-114}. In the epoxidation of (II) the *trans*-double bond reacts first with formation of the epoxydiene (XIV)⁶⁸:



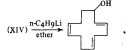
It has been established ¹¹² that monoepoxidation of (II) results in the formation of 92% of the *trans*-epoxide and 7% of the *cis*-epoxide. The epoxidation of (I) yielded the epoxydiene (XV). The oxidation of (I) and (II) by atmospheric oxygen in the presence of boric acid or its esters leads to the formation of epoxycyclododecadiene together with 2,6,10-cyclododecatrienol ¹¹⁵. The synthesis of 5,6:9,10-diepoxycyclododecene, 5,6:9,10-diepoxycyclododecane ¹¹⁶, 1,2:5,6:9,10-triepoxycyclododecane ¹¹⁷, and epoxycyclododecene ^{118,119} has been described in patents. The opening of the epoxy-rings of these compounds results in the formation of the corresponding diols, tetraols, and hexaols ^{68,74,120-122}; thus (XIV) and (XV) give rise to the corresponding *trans*-dihydroxycyclododecadienes ⁶⁸. The reduction of double bonds in (XIV) and (XV) leads to *trans*-epoxycyclododecane and subsequent opening of the epoxy-ring gives rise to *cis*-1,2-cyclododecaneil0^{68,123}:

$$(X1V) \quad \text{or} \quad (XV) \xrightarrow{H_2} (CH_2)_{10} \stackrel{CH}{\underset{CH}{\longrightarrow}} 0 \xrightarrow{H^{\bigoplus}}_{H_20} (CH_2)_{10} \stackrel{CH-OH}{\underset{CH}{\longrightarrow}} H^{\bigoplus}$$

In the presence of catalytic amounts of Group I and II metal halides epoxycyclododecadienes readily isomerise to the corresponding ketones 124, 123:

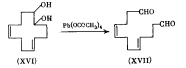


Two isomeric unsaturated ketones with different positions of the keto-group in relation to the C=C double bonds are formed from (XIV).¹²⁵ When (XIV) is acted upon by (iso- $C_4H_9)_2AlH$, ¹²⁶ n- C_4H_9Li , or C_4H_9MgBr ¹³⁷ isomerisation with formation of unsaturated alcohols takes place:

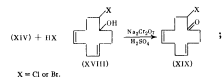


The isomerisation of the epoxide and the formation of cyclodode canone 11,75,110 together with cyclodode canol are observed in the catalytic hydrogenation of (XIV) or (XV) in the presence of nickel catalysts (Raney nickel, Ni/Al₂O₃, Ni/Cr₂O₃) at 120-150°C.

When (II) is oxidised by OsO₄ or KMnO₄, the reaction proceeds primarily at the *trans*-double bond with formation of the corresponding *trans*-diol and *trans*, *trans*tetraol¹²⁸. When cyclododecadienediol (XVI) was oxidised, the unsaturated dialdehyde (XVII) was obtained¹²⁹:

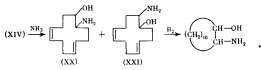


The opening of the epoxy-ring in (XIV) and (XV) takes place readily under the influence of hydrochloric and hydrobromic acids with formation of the corresponding halogenohydrins $(XVIII)^{114}$:

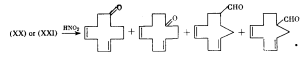


When the halogenohydrins (XVIII) are oxidised by the chromic acid mixture, it is possible to obtain high yields of the unsaturated halogeno-ketones (XIX). The halogenohydrins and the halogeno-ketones obtained from the epoxide (XIV) consist of a mixture of isomers with different positions of the OH group, the halogen atom, and the carboxy-group relative to the *cis*- and *trans*-double bonds in the ring ¹¹⁴. Cycloundecadienecarboxylic acid has been obtained from the ketones (XIX) by the Favorskii reaction ¹¹⁴:

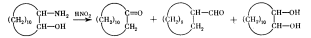
On heating with an aqueous ammonia solution, the epoxide (XIV) gives rise to a mixture of two isomeric *cis*-aminoalcohols (XX) and (XXI)¹³⁰, which form *cis*-1,2-aminocyclodode canol on hydrogenation:



When (XX) and (XXI) were deaminated, a mixture of isomeric cyclododecadienones and formylcycloundecadienes was isolated ¹³⁰:

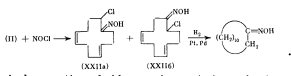


Regardless of the configuration of the initial compound, the deamination of cis- and trans-2-aminocyclododecanols leads to a mixture of cyclododecanone, formylcycloundecane and cis-1,2-cyclododecanediol in equal proportions ¹³⁰:



When α -aminocyclodode canecarboxylic acid is deaminated, α -hydroxycyclodode canecarboxylic acid is obtained in a quantitative yield¹³¹.

Nitrosyl chloride readily adds to (II) at a *trans*-double bond with formation of two products (XXII), a and b)¹³²⁻¹⁴⁰:

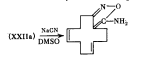


The hydrogenation of chloro-oximes at atmospheric pressure in the presence of platinum and palladium catalysts yields cyclododecanone oxime as the main product ¹³², ^{133,135}. Several patents ¹⁴¹⁻¹⁴⁴ and another communication ¹⁴⁰ describe chlorine exchange reactions involving α chlorocyclododecadienone oxime. When (XXIIa) and (XXIIb) are treated with nucleophiles, products resulting from the substitution of a chlorine atom are obtained in a high yield ¹⁴²:

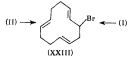
$$(XXIIa) + (XXII6) \xrightarrow{X\Theta} +$$

 $X = C_2H_5O$, CH_3O , C_2H_5S , CH_3COO , or R_2N .

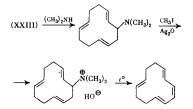
The reaction of (XXIIa) with sodium cyanide in dimethyl sulphoxide (DMSO) leads to the formation of the corresponding aminoisoxazoles ¹⁴⁰, for example:



Allylic bromination of (II) by *N*-brom osuccinimide proceeds with isomerisation involving the *cis*-double bond and leads to bromocyclododecatriene (XXIII), which is also obtained on allylic bromination of (I)⁵⁸:

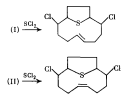


Treatment of the bromo-derivative (XXIII) with ammonia, dimethylamine, and potassium acetate gives rise to substitution products⁵⁸, for example:

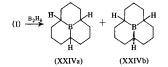


A cylation of (II) via the Kondakov reaction yields a mixture of 1-acyl-2,5-9- and 1-acyl-1,5,9-cyclododecatrienes¹⁴⁵. Reactions involving the addition of sulphur

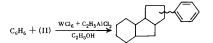
dichloride to (I) and (II) and leading to dichlorothiabicyclotridecenes are of interest 146 , 152 :



The chlorine atoms in these compounds are readily replaced by cyano- and acetoxy-groups ¹⁵². The hydroboration of (II) gives rise to a mixture of the *cis*- and *trans*isomers of 9*b*-boraperhydrophenalene (XXIV, a and b) together with 9*b*-boraperhydrobenzo[*ed*]azulene. The latter is converted into *cis*-9*b*-boraperhydrophenalene on heating ¹⁴⁷, ¹⁴⁸:



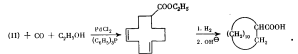
(XXIVa) readily reacts with LiH to form an addition product, which is an effective stereoselective reducing agent for cyclic ketones ^{149,150}. Trichlorosilane and dichloromethylsilane add to (II) preferentially at the *trans*double bond in the presence of platinum ¹⁵¹. When (II) is treated with hydrogen bromide in the presence of benzoyl peroxide, 9-halogeno-1,5-cyclododecadiene is obtained ¹⁵⁴. The reaction of (II) with nitrogen oxides has been described ¹⁵⁵. When (II) reacts with benzene in the presence of the WCl₆ + C₂H₅AlCl₂ + C₂H₅OH system, phenyltricyclo[6,4,0,0^{2,6}]dodecane is formed ¹⁵⁶:



In complete ozonisation of (I) and (II) with subsequent cleavage of the ozonide leads to deca-3,7-diene-1,10-dicarboxylic acid and the corresponding dialdehyde 157,158 :

(I) or (II) $\xrightarrow{O_3}$ HOOC (CH₂)₂ CH=CH (CH₂)₂ CH=CH (CH₂)₂ COOH .

It has been established that ozone attacks (I) and (II) at the same rate¹⁵⁹. The carbonylation of (I) and (II) under the influence of CO in the presence of metal (Ni, Co) carbonyls and palladium complexes with formation of cyclododecane-carboxylic acid has been described in several communications¹⁶⁰⁻¹⁶⁷. The carbonylation of (II) with the aid of palladium catalysts in alcohol gives rise to a high yield of . ethyl 4,8-cyclododecadienecarboxylate¹⁶²:



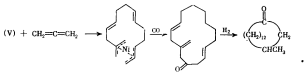
When cyclododecanecarboxylic acid is acted upon by nitrosylsulphuric acid, ω -dodecalactam is obtained in a high yield ^{168,169}:



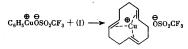
Certain other reactions of cyclododecanecarboxylic acid have been described 170 , (I) and (II) react on heating with maleic anhydride to form 1:1 adducts $^{172-174}$. The

condensation of (II) with cyclopentadiene has been described $^{\rm 175}.$

Metal complexes of (I) and (II) are known. A π -allyl complex of (V), which reacts with allene, has been described ¹⁷⁶,¹⁷⁷. Subsequent reactions of the product lead to the synthesis of DL-muscone:



 π -Complexes of palladium $(C_{12}H_{17-19}PdCl)_2^{178}$ and copper ¹⁷⁹, ¹⁸⁰ have been obtained and a copper complex, synthesised by the reaction ^{181,182}



has been synthesised.

When a heptane solution of (I) is refluxed with bis(tri-methylgermyl)tetracarbonylruthenium, a mixture of products with different structures is formed¹⁸³. The reaction of (I) with dode cacarbonyltriruthenium also leads to a mixture of products having different compositions¹⁸⁴:

$$\begin{array}{l} ({\rm I}) + {\rm Ru}_3\,({\rm CO})_{12} \ \rightarrow \ {\rm HRu}_3\,({\rm CO})_{9}\,({\rm C}_{12}{\rm H}_{16}) + {\rm HRu}_3\,({\rm CO})_{9}\,({\rm C}_{12}{\rm H}_{17}) + \\ \\ + \ {\rm HRu}\,({\rm CO})_{7}\,({\rm C}_{23}{\rm H}_{34}) + {\rm Ru}_4\,({\rm CO})_{10}\,({\rm C}_{12}{\rm H}_{16}) \end{array}$$

Analogous complexes have been obtained for rhodium 185,186 , iridium 187 , nickel 40,48 , and iron 188 .

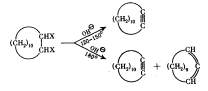
Silver nitrate forms with (I) three types of adducts in which one, two, or three *trans*-double bonds are coordinated to the silver ion 56,68,189:

 $C_{12}H_{18} \cdot AgNO_3, \quad C_{12}H_{18} \cdot 2AgNO_3, \quad C_{12}H_{18} \cdot 3AgNO_3.$

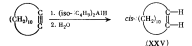
When (II) reacts with a silver ion, only the two *trans*double bonds are coordinated, while the *cis*-double bond is unaffected 68,189 .

IV. SYNTHESES BASED ON *cis*- AND *trans*-CYCLODODE-CENES

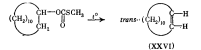
Several methods for the synthesis of *cis*- and *trans*cyclododecenes (XXV) and (XXVI) are known. It was noted above that they are obtained on selective hydrogenation of (I) and (II)^{83,964,91-94} in the presence of homogeneous catalysts or when (II) is acted upon by di-imide⁸⁰. A mixture of (XXV) and (XXVI) is formed when HCl is eliminated from chlorocyclododecane in the presence of iron-containing catalysts¹⁹⁰. Dehalogenation of 1,2-dichloro- or 1,2dibrom o-cyclododecane by sodionaphthalene or sodiobiphenyl gives a 90% yield of a mixture of (XXV) and (XXVI)¹⁹¹. When 1,2-dihalogenocyclododecanes are heated in the presence of alkalis at 150°C, only cyclododecyne is formed¹⁹²⁻¹⁹⁴. On raising the temperature to 180°C, a mixture of 28% of 1,2-cyclododecalene and 72% of cyclododecyne was isolated¹⁸³:



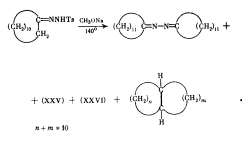
An analogous mixture was obtained after double dehydrobromination of dibromocyclododecane with alcoholic alkali ¹³⁰. When cyclododecyne is reduced by $(iso-C_4H_9)_2$. .AlH, mainly 6-cyclododecene is formed ¹⁹³:



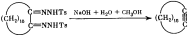
A mixture of 90% of *cis*- and 10% of *trans*-cyclododecyne is formed on decomposition of cyclododecenone semicarbazone by alkoxides at 180–240°C.^{195,196} Pyrolysis of cyclododecyl acetate at 550°C leads to a mixture of isomers (XXV) and (XXVI) in proportions of 1:5.¹⁹⁷ When (XXV) or (XXVI) is heated in acetic acid at 80–130°C, *cis*-*trans* isomerisation takes place with formation of an equilibrium mixture containing approximately equal amounts of (XXV) and (XXVI)¹⁹⁷. Contrary to the Chugaev rule, the decomposition of *O*-cyclododecyl *S*methyl thiocarbonate gives rise to *trans*-cyclododecene¹³⁰:



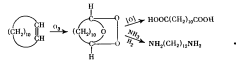
When cyclododecanol was heated to $250-270^{\circ}$ C in the presence of toluene-*p*-sulphonic acid, a high yield of cyclododecene was obtained ¹⁹⁸. The reaction of cyclododecene with *N*-bromosuccinimide and subsequent dehydrobromination of the product in pyridine or quinoline result in the formation of 1,3-cyclododecadiene¹⁹⁹. The dehydration of 6-hydroxy- or 6-acetoxy-cyclododecanone in the presence of Zn₃(PO₄)₂ and Fe₂O₃, leading to cyclododec-5-en-1-one, has been described. In the presence of alkyl-lithium in tetrahydrofuran (THF), cyclododecanone *N*, *N*-ditosylhydrazone decomposes to form a mixture of *cis*- and *trans*-cyclododecene and cyclododecanone²⁰¹. When cyclododecanone tosylhydrazone is heated in the presence of sodium methoxide, the decomposition leads mainly (to the extent of 60%) to cyclododecanone azine²⁰²:



On photolysis of 1,2-cyclododecanedione bistosylhydrazone in an alkaline solution, cyclododecyne was isolated in 53% yield ²⁰³:



Ozonisation with subsequent conversion into decane-1,10dicarboxylic acid²⁰⁴⁻²⁰⁶ and 1,12-diaminododecane²⁰⁷ is an important reaction of cyclododecene:



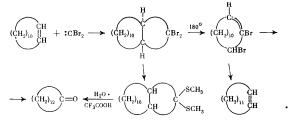
When cyclododecene is oxidised by nitric acid in the presence of ammonium vanadate, considerable amounts of nonane-1,9-dicarboxylic acid are formed together with decane-1,10-dicarboxylic acid 208 . Catalytic oxidation of cyclododecene by hydrogen peroxide in the presence of Re₂O₇ takes place with dissociation of the C=C bond and yields decane-1,10-dicarboxylic acid 209 . Oxidation with potassium permanganate in acetic anhydride leads to 1,2cyclododecanedione (48%) and other products 210 :

In an alkaline medium in the presence of catalytic amounts of benzyltriethylammonium chloride, trans-1,2-cyclodode-canediol is formed in 50% yield²¹¹.

Cyclododecanecarboxylic acid has been obtained from cyclododecene by the Koch reaction ¹⁶⁷:

$$(CH_2)_{10}$$
 $\overset{CH}{\underset{GH}{\parallel}}$ $\overset{CO}{\underset{BF_3}{\leftarrow}}$ $(CH_2)_{10}$ $\overset{CHCOOH}{\underset{CH_2}{\leftarrow}}$

Dibrom ocarbene adds smoothly to cyclododecenes with formation of 13,13-dibrom obicyclo[10,1,0]tridecane, from which cyclotridecanone^{212,213} and cyclotridecene²¹² were obtained:

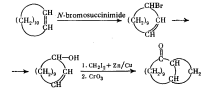


Metathesis of cyclododecene in the presence of the WCl₅ + C₅H₅AlCl₂ system yielded unsaturated cyclic hydrocarbons containing 24,36, and 48 carbon atoms in the ring²¹⁴. The condensation of cyclododecene and (II) with formaldehyde has been described ²¹⁵:

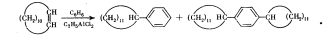
$$(CH_2)_{10} \parallel H + HCHO + (CH_3CO)_2O \longrightarrow (CH_2)_{10} \parallel CH = CH_2OCOCH_3$$

When cyclododecene reacts with paraformaldehyde, transbicyclo[10,3,0]-2-oxa-14-pentadecene is formed ²¹⁶.

trans-Bicyclo[10,1,0]-2-tridecanone has been obtained from cyclodode cene²¹⁷:



When cyclododecene is acted upon by NO in the presence of O₂, 1-nitroso-2-nitrocyclododecane is obtained. It rearranges on heating in DMF to give 2-nitrocyclododecanone oxime. 1,12-Diaminododecane and ω -nitrolauric acid were obtained from the latter ²¹⁸. Cyclododecene undergoes a Friedel-Crafts reaction with benzene and its homologues to form alkylation products ^{219,220}.



Pyrolysis of (XXV) and (XXVI) results in the cleavage of the ring with formation of 1,8-nonadiene and 1,11-dodecadiene and also in isomerisation involving the double bonds²²¹. When cyclododecene was treated with atmospheric oxygen in t-C₄H₉OH in the presence of manganese naphthanate, 1,2,3-cyclododecanetriol was obtained²²². Photochemical cycloaddition of the olefins (XXV) and (XXVI) to dichlorovinylene carbonate leads to three stereoisomeric carbonates of dichlorobicyclo[10,2,0]-2,3-tetradecanediols²²³.

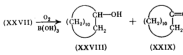
The synthesis of ω -cyanoundecanoic aldehyde from (XXV) and (XXVI) via the mechanism

$$(\underbrace{CH_{2^{1}l_{0}}}_{CH} \underbrace{\overset{CH}{\underset{H}{\longrightarrow}}}_{CH} \underbrace{(\underbrace{CH_{2^{1}l_{0}}}_{CHCL}}_{CHCL} \underbrace{\overset{CH}{\underset{NaOH}{\longrightarrow}}}_{AOH} \underbrace{(\underbrace{CH_{2^{1}l_{0}}}_{CHOCH_{3}} \underbrace{\overset{PCl_{5}}{\underset{H}{\longrightarrow}}}_{NC(CH_{2})_{l_{0}}CHO} NC(CH_{2})_{l_{0}}CHO$$

is of great interest ²²⁴. The reaction of cyclododecene with B_2H_6 yielded the corresponding tri(cyclodecyl)borane ²²⁵. When cyclododecene was treated with nitrogen oxides and air, α -nitrocyclododecanone was obtained in a high yield ²²⁵⁻²³⁰. The mechanism of the oxidation of (XXV) and (XXVI) by molybdenum peroxide has been investigated ²³¹. A study has been made of the kinetics of the hydrogenation of (XXV) and (XXVI) to cyclododecane in the presence of platinum and palladium blacks ²³². (XXV) and (XXVI) form π -allyl complexes with palladium chloride ²³³. The structure of the complex of *trans*-cyclododecene with AgNO₃, having the composition (*trans*-C₁₂H₂₂)₂.AgNO₃, has been investigated ²³⁴. On heating with perfluorobutyne, *cis*, *trans*-1,3-cyclododecadiene gives rise to a mixture of two bicyclic products ²³⁵.

V. SYNTHESES OF CYCLODODE CANE DERIVATIVES

The methods of synthesis of cyclododecane derivatives are of great significance, since the preparation of the industrially important decane-1,1-dicarboxylic acid, 1,12diaminododecane, and ω -dodecalactam is based on them. The oxidation of cyclododecane (XXVII) by atmospheric oxygen to cyclododecanol (XXVIII) and cyclododecanone (XXIX) has been studied in detail. When (XXVII) is oxidised by oxygen in the presence of boric acid or boron trioxide, a mixture of 80% of (XXVIII), 8-10% of (XXIX), and 10% of polyfunctional oxygen-containing compounds (conversion 30-35%) is formed ^{69,236}:



The high selectivity of the oxidation in the presence of boric acid has been pointed out in patents $^{237-239}$. A greater amount of the ketone (XXIX) is formed in the presence of manganese or cobalt salts 240,241 . When (XXVII) was oxidised by oxygen in the absence of boric acid, it was found that at the beginning of the process cyclododecyl hydroperoxide, (XXIX), and (XXVIII) accumulate simultaneously $^{242-245}$. After the attainment of the maximum, the concentration of the hydroperoxide falls owing to its thermal decomposition. This results in the formation of a mixture of 70% of (XXIX) and 30% of (XXVIII). Cyclododecane is oxidised to cyclododecyl hydroperoxide in the presence of alkali metal salts $^{246-248}$. A study of the mechanism of the decomposition of cyclododecyl hydroperoxide showed 249 that considerable amounts of undecane and undecyl alcohol are formed in an inert medium, while in the presence of O₂ hardly any undecane is formed. The oxidation of (XXVII) by nitrogen dioxide to decane-1,10-dicarboxylic acid has been reported 250,251.

Treatment of (XXVII) by hydroxylamine and oxygen under irradiation leads to 1,2-cyclododecanedione monoxime²⁵²:

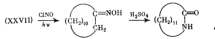
(XXVII)
$$\xrightarrow{\text{NH}_2\text{OH} + 0}_{\text{CuCl}_2, hy}$$
 (CH₂)₁₀ C=NOH

Nitrocyclododecane was obtained in the liquid-phase nitration of (XXVII) by nitric acid ^{168,253,254}:

$$(XXVII) \xrightarrow{HNO_3} (CH_2)_{10} \xrightarrow{CH-NO_2} CH_2$$

Nitrocyclododecane was also obtained on hydrogenation of 3-nitrocyclododecene²⁵⁵. When nitrocyclododecane is treated with a solution of potassium hydroxide, the potassium salt of *aci*-nitrocyclododecane is formed and is converted into *aci*-nitrocyclododecane on acidification²⁵⁴,²⁵⁶. This *aci*-form is stable and does not change even on recrystallisation²⁵⁴. When nitrocyclododecane was hydrogenated, aminocyclododecane and cyclododecanone oxime were obtained²⁵⁴,²⁵⁷. When the potassium salt of *aci*-nitrocyclododecane was treated with H₂NOH.HCl, cyclododecane oxime is formed in a high yield²⁵⁴.

The hydrolysis of a nitrocyclododecane salt results in the formation of cyclododecanone $^{254},^{258},^{259}$. The conversion of nitrocyclododecane into ω -dodecalactam at a high temperature in the presence of certain metal oxides has been described 260 . Numerous studies have been devoted to the photonitrosation of (XXVII) to cyclododecanone oxime $^{139},^{153},^{168},^{261-266}$ and to the isomerisation of cyclododecanone oxime to ω -dodecalactam, which is the initial compound in the synthesis of polyamide-12 $^{261},^{262},^{267-271}$:

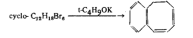


Nitrodode calactam has been obtained by treating α -nitrocyclodode canone oxime with concentrated sulphuric acid at 40-50° C:²⁷²



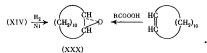
The acid or alkaline hydrolysis of ω -dodecalactam results in the formation of ω -aminododecanoic acid^{11,273},274</sup>.

The photochemical chlorination of (XXVII) to monochlorocyclododecane has been investigated ^{75,275,276}. Monochlorocyclododecane undergoes the Friedel-Crafts reaction in benzene and toluene, forming phenyl- and tolyl-cyclododecanes ²⁷⁷. Hexabromocyclododecane, obtained from (I) and (II) on dehydrobromination, undergoes an unusual transformation into benzocyclo-octatriene ²⁷⁸:



When (XXVII) is heated to 300°C in the presence of platinised charcoal, it undergoes transannular dehydrocyclisation with formation of bicyclic hydrocarbons²⁷⁹.

Epoxy cyclodode cane (XXX) has been obtained by the hydrogenation of the double bonds in epoxy cyclodode cadiene 68 and by the epoxidisation of cyclodode cene by peracids 75,130 :



The *cis*- and *trans*-oxides are formed from *cis*- and *trans*cyclododecene. Catalytic hydrogenation of (XXX) over Raney nickel^{11,68,75} leads to the formation of cyclododecanol and cyclododecanone:

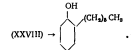
$$XXX) \xrightarrow{H_{\bullet}} (XXVIII) + (XXIX)$$

Isomerisation of the epoxide (XXX) under the influence of LiBr^{125} or MgL_2^{124} leads to cyclododecanone:

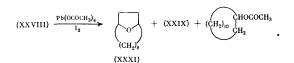
$$(XXX) \xrightarrow{MgI_2} \rightarrow (XXIX) \cdot$$

Cyclododecanol has been obtained by hydrating cyclododecene with sulphuric acid ²⁸⁰ and also on hydrogenation and hydrolysis of 1-acetoxy-5,9-*trans*, *trans*-cyclododecadiene²⁷³. Dehydrogenation of cyclododecanol at 240-250°C under the influence of catalysts (Raney nickel, Cu/Cr, Ni/Al₂O₃, Cu/SiO₂) leads to the formation of cyclododecanone in a high yield^{11,70},²⁴⁰,²⁸¹. The synthesis of cyclododecanone by the hydrolysis of cyclododecanone oxime has been described^{258,259}.

Decane-1,10-dicarboxylic acid is formed in a high yield when cyclododecanol or its mixture with cyclododecanone is acted upon by nitric acid in the presence of ammonium metavanadate ^{11,110,282} or V₂O₅.²⁸³ It has been stated in patents ^{284,285} that the oxidation of cyclododecanol is accompanied by nitration, which resulted in the isolation of ω , ω -dinitrododecanoic acid; however, the addition of copper powder to the oxidation reaction prevents the formation of nitro-acids ²⁸⁵. When cyclododecanol is treated with a mixture of FSO₃H-SbF₅-SO₂, it isomerises to hexylcyclohexanol²⁸⁶:

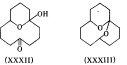


The oxidation of cyclododecanol by lead tetra-acetate in the presence of iodine gives rise to a mixture of oxygencontaining products in which 13-oxabicyclo[8,2,1]dodecane (XXXI) predominates 287 :



Oxidation in the absence of iodine leads mainly to acetoxycyclododecane.

When cyclododecan ediol is heated in the presence of Raney nickel, 2-hydroxycyclododecanone is formed ²⁸⁸, while catalytic oxidation in the presence of $[(C_6H_5)_3P]_3RuCl_2$ and benzaldehyde yields 1,2-cyclododecanedione ²⁸⁹. Methods for the preparation of 2-alkoxy-1-alkylcyclododecanes ²⁹⁰ and cyclododecylamides ²⁰¹, which exhibit pharmacological activity, have been described. 1-(Dialkylaminoalkyl)cyclododecyl carboxylates ²⁹² and the esters of 1-(aminoalkyl)cyclododecanol possess bactericidal properties²⁹³. The oxidation of *cis*, *cis*, *trans*-1,5,9-cyclododecanetriol by the Brown method proceeds with a transannular interaction and leads to the hemiacetal (XXXII) and the acetal (XXXIII) in proportions of 43:57.²⁹⁴

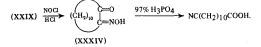


VI. SYNTHESES BASED ON CYCLODODE CANONE

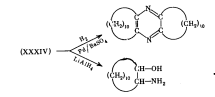
Numerous compounds have been synthesised on the basis of cyclododecanone (XXIX) and many interesting reactions have been carried out. As stated above, (XXIX) is obtained by the dehydrogenation of cyclododecanol (XXVIII)^{11,70,240,281}, the isomerisation of epoxycyclododecanol (XXVIII)^{124,125}, the hydrolysis of cyclododecanone oxime ^{258,259}, and the oxidation of cyclododecane^{69,236-248} and cyclododecanol⁶⁹. Several studies have been devoted to the synthesis of cyclododecanone oxime from (XXIX) and its conversion into ω -dodecalactam ^{11,70,295-297}.

The reaction of hydroxylamine with cyclododecanone under the influence of hydroxylammonium chloride or sulphate takes place readily and in a quantitative yield in the presence of bases (ammonia, potassium or sodium carbonate)^{11,70,295,296}. When industrial hydroxylammonium sulphate, containing ammonium sulphate and sodium sulphate, is used, the conversion of cyclododecanone into the oxime takes place without the addition of bases, because sulphuric acid is bound into acid ammonium and sodium salts^{11,295}. The reaction with hydroxylamine is carried out in alcoholic, aqueous alcoholic, and aqueous media, a higher temperature being necessary in an aqueous medium. Cyclododecanone oxime forms a 2:1 complex with SnCl4.²⁹⁸

The mechanism of the Beckmann rearrangement in cyclododecanone oxime has been investigated and it has been found that the process proceeds via a unimolecular mechanism²⁹⁹. α -Chlorocyclododecanone oxime is readily obtained when nitrosyl chloride adds to cyclododecene³⁰⁰. The substitution of the chlorine atom in α -chlorocyclodo-decanone oxime by alkoxy- and amino-groups has been described³⁰¹⁻³⁰⁴. On treatment of (XXIX) with NOCl or RONO₂ in the presence of HCl or sodium alkoxide, cyclo-dodecanedione monoxime (XXXIV) is formed in a high yield³⁰⁵; it rearranges smoothly to 11-cyanoundecanoic acid under the influence of phosphoric acid^{306,307}:



The hydrogenation of (XXXIV) yielded bis-2,3,5,6-deca-methylenepyrazine 130 :



The reaction of (XXIX) with NOCl and ethanol in SO₂ leads to the cleavage of the ring via the mechanism ³⁰⁸:

$$(XXIX) + NOCI + C_2H_5OH \xrightarrow{SO_2} \rightarrow HON = CH (CH_2)_{10} COOC_2H_5$$

 α -Nitrocyclododecanone is obtained by the hydrolysis of its oxime with hydrochloric acid³⁰⁹. The alcoholysis of α -nitrocyclododecanone results in the formation of ω nitrododecanoate esters³¹⁰:

When (XXIX) is brominated, monobromo- or dibromocyclododecanone³¹¹ is obtained:

(XXIX)
$$\xrightarrow{\text{Br}_2}$$
 (CII₂)₁₀ $\xrightarrow{\text{CHBr}}$ $\xrightarrow{\text{Br}_2}$ (CII₂)₁₀ $\xrightarrow{\text{CHBr}}$ $\xrightarrow{\text{CI}_4}$ (CII₂)₉ C=0 CIIBr

Monohalogenocyclododecanones are readily formed on oxidation of the corresponding halogenohydrins¹¹⁴:

$$(\overset{CH_{2})_{10}}{\underset{CH0H}{\overset{CHX}{\longrightarrow}}} \overset{Cr0_{3}}{\underset{CH0}{\longrightarrow}} (\overset{CH_{2})_{10}}{\underset{CH0}{\overset{CHX}{\longrightarrow}}} \overset{CHX}{\underset{CH2}{\longrightarrow}} ;$$

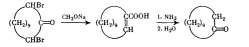
X = Cl, or Br.

The Favorskii rearrangement of α -halogenocyclododecanones under the influence of alcoholic alkali leads to the formation of cycloundecanecarboxylic acid¹¹⁴:

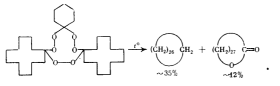
$$(CH_{2^{1}l_0} CHX \longrightarrow (CH_{2^{1}l_0} CHCOOH ;$$

 $X = Cl, \text{ or Br.}$

Under these conditions, α , α' -dibromocyclododecanone gives rise to cycloundecenecarboxylic acid, which has been converted into cycloundecanone^{311,312}:



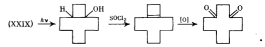
The oxidation of (XXIX) by hydrogen peroxide in the presence of SeO₂ leads to cycloundecanecarboxylic acid³¹³. Cyclododecanone peroxide has been obtained by treating (XXIX) dissolved in carboxylic acids with H_2O_2 .^{314,315} In the presence of cyclic ketones bis(cyclododecylidene) cycloalkylidene triperoxides are formed³¹⁶ and their thermal decomposition in chlorobenzene leads to macrocyclic compounds, for example:



Perbenzoic acid $^{317-319}$, Caro's acid, and hydrogen peroxide in sulphuric acid 318 were used to oxidise (XXIX) to dodecanolide:



Irradiation of (XXIX) results in a transannular rearrangement with formation of $bicyclo[8,2,0]dodecan-1-ol^{320}$,



On heating cyclodode canone and cyclodode canol with alkalis to 350° C, a mixture of lauric and nonane-1,9-dicarboxylic acid is formed in 90% yield ³²²⁻³²⁴:

 $(XXVIII) + (XXIX) \xrightarrow{KOH} CH_3 (CH_2)_{10} COOH + HOOC (CH_2)_3 COOH$.

The addition of hydrocyanic acid to (XXIX) takes place readily in the presence of $(C_2H_5)_{3N}$ with formation of a cyanohydrin $^{325-327}$, from which cyclododecanecarboxylic acid 326 , cyclotridecanone 327 , and brassylic acid 326 were obtained:



The formation of hydantoins from cyclododecanone or cyclododecenones has been described ³²⁸:

$$(XXIX) + KCN + (NH_4)_2CO_3 \longrightarrow (CH_2)_1 C NHCO CONH$$

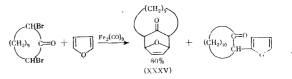
1,2-Cyclododecanedione is formed in a high yield on heating hydroxycyclododecanone with copper diacetate³²⁹, $\frac{330}{2}$:

$$(\underbrace{\operatorname{CH}_2}_{l_0})_{l_0} \xrightarrow{\operatorname{CHB}_r} \xrightarrow{\operatorname{OH}_{\Theta}} (\underbrace{\operatorname{CH}_2}_{l_0})_{l_0} \xrightarrow{\operatorname{CH}_{\Theta}} (\underbrace{CH}_2}_{l_0})_{l_0} \xrightarrow{\operatorname{CH}_{\Theta}} (\underbrace{CH}_2}_{l_0})_{l_0} \xrightarrow{CH}_{O} (\operatorname{CH}_2}_{l_0})_{l_0} \xrightarrow{CH}_{O} (\operatorname{CH}_2}_{l_0})_{l_0} (\underbrace{CH}_2}_{l_0})_{l_0} \xrightarrow{CH}_{O} (\operatorname{CH}_2}_{l_0})_{l_0} (\underbrace{CH}_2}_{l_0})_{l_0} (\operatorname{C$$

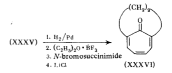
Diazocyclododecanone has been obtained by the reaction³³¹:

$$(CH_2)_{10} \xrightarrow{C=0} (CH_2)_{10} \xrightarrow{CH_3C_6H_4So_2N_3} (CH_2)_{10} \xrightarrow{C=0} (CH_2)_{10} \xrightarrow{C=0} N_2$$

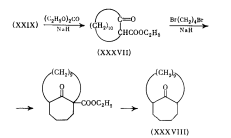
 α -Halogenocyclodode canones are readily reduced to cyclodode canone in the presence of LiI and $(C_2H_5)_2O.BF_3$.³³² The reaction of α , α' -dibromocyclodode canone with furan in the presence of Fe₂(CO)₉ takes place in an interesting manner³³³:



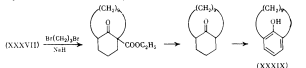
(XXXV) has been converted via a series of stages into tropone (XXXVI) 334 :



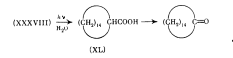
Tropone has been obtained from cyclododecanone also in accordance with another scheme via the ethoxycarbonyl derivative $(XXXVII)^{335}$:



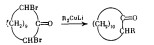
When the ketone (XXXVIII) was brominated and debrominated, tropone was obtained 336 . Metacyclophane (XXXIX) was synthesised similarly 336 :



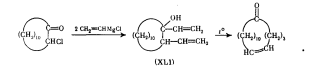
Ultraviolet irradiation of the ketone (XXXVIII) led to the formation of cyclopentadecanecarboxylic acid (XL), which was converted into cyclopentadecanone 335,337 :



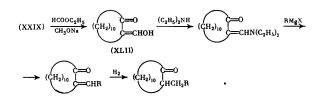
A new method has been proposed for the alkylation of α , α' -dibromocyclodode canone with formation of α -alkyl-cyclodode canones^{338,339}:



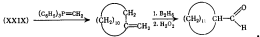
The action of a Grignard reagent on (XXIX) with formation of alkylcyclododecanols has been investigated 202 . Methylcyclododecanone has been obtained by the reaction of bromocyclododecanone with CH₃ZnI in DMSO 202 . The reaction of chlorocyclododecanone with two moles of vinylmagnesium chloride leads to the formation of (XLI), which is converted into 5-cyclohexadecenone after the Cope rearrangement 340 , 341 :



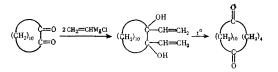
 α -Alkylcyclododecanones have been obtained by the formylation of (XXIX) via the mechanism³⁴²:



(XLII) undergoes oxidative cleavage by hydrogen peroxide, which leads to brassylic acid³²⁵. Formylcyclododecane was synthesised from (XXIX) as follows²⁰²:



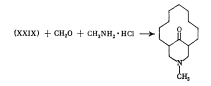
A method has been described for the synthesis of 1,6cyclohexade canedione by treating 1,2-cyclodode canedione with vinylmagnesium chloride and a subsequent rearrangement³⁴³:



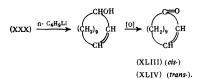
Depending on temperature, the isomerisation of 2,3-epoxycyclododecanone takes place with formation of 1,2-or 1,3-cyclododecanediones³⁴⁵:



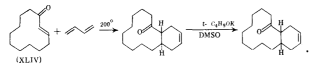
Methods have been developed for the synthesis of [7]metacyclophane and its bromo-derivative via the aldol condensation of 1,4-cyclododecanedione 346 , 347 and [7](2,6)pyridinophane derived from 1,5-cyclododecanedione 348 , 349 . Cyclododecanone has been made to undergo the Mannich reaction 350 , 351 :



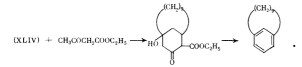
The *cis*- and *trans*-2-cyclododecenones (XLIII) and (XLIV) were obtained from epoxycyclododecane³⁵²:



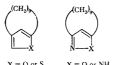
The synthesis of a mixture of cis- and trans-cyclododecenones by the bromination of cyclododecanone ethylene acetal and subsequent dehydrobromination by t-C₄H₉OK in DMSO has been proposed³⁵³. cis-Cyclododecenone (XLIII) is thermodynamically less stable than the trans-isomer (XLIV) and isomerises to the latter³⁵². Ultraviolet irradiation of a hexane solution of (XLIV) leads to a mixture of (XLIII) and cis-3-cyclododecenone³⁵². Diene condensation of (XLIV) with butadiene leads to trans-bicyclo[10,4,0]hexadec-14(15)-en-2-one, which wholly isomerises to the cis-isomer on treatment with t-C₄H₉OK:³⁵⁴



The cis-isomer (XLIII) does not undergo diene condensation with butadiene. [9]Metacyclophane and [9](2,4)pyridinophane have been obtained by the reaction of (XLIV) with a cetoacetic or cyanoacetic ester and subsequent reac-tions of the adducts³⁵⁵:

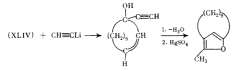


A series of [9] heterophanes have been obtained from (XLIV)^{356,357}:

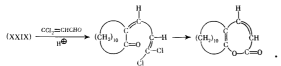


X = O or SX = O or NH

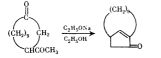
When (XLIV) reacts with hydrazine hydrate and this is followed by dehydrogenation with sulphur, [9](3,5) pyrazolophane is formed, while treatment with lithium acetylide and further cyclisation lead to 11-methyl[9](2,4)furano-phane³⁵³:



Decamethylene- α -pyrone has been obtained by the condensation of (XXIX) with dichloroacrolein and subsequent cyclisation 358:

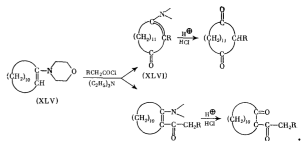


The product of the condensation of 1-methoxycyclododecene with 2-methylbut-3-yne-2-ol undergoes the Cope rearrangement to give 3,3-dimethylallenylcyclododecenone, which isomerises on ultraviolet irradiation to 4-isopropylidenecyclotetradecenone³⁵⁹. When 3-acetylcyclododecan one is acted upon by C_2H_5ONa , a bicyclic ketone is formed ³⁶⁰:

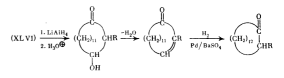


Enamines have found extensive applications in the synthesis of various cyclodode canone derivatives $^{361-363}$. When morpholinocyclododecene (XLV) is treated with carboxylic acid chlorides in the presence of $(C_2H_5)_3N$, 2-alkyl-1,3cyclotetradecanediones (resulting from the insertion of two

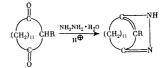
carbon atoms in the ring) and acylation products are formed 364,365.



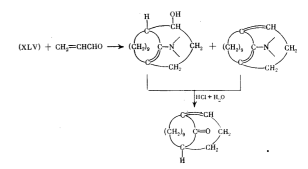
When (XLVI) was treated with lithium tetrahydroaluminate and this was followed by dehydration and hydrogenation, methyl- and ethyl-cyclotetradecanones, which have a strong musk-like scent, were obtained ³⁶⁶:



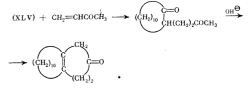
 $R = CH_3, C_2H_5$ The interaction of 2-alkyl-1,3-cyclotetradecanedione with hydrazine hydrate³⁶⁴ or 2,4-dinitrophenylhydrazine³⁶⁶ leads to a pyrazole derivative:



When acrolein acts on (XLV), an interesting reaction takes place³⁴⁴:



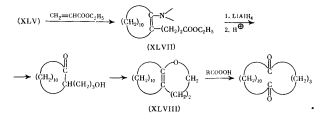
The interaction of (XLV) with methyl vinyl ketone leads to a bicyclic ketone³⁶⁷:



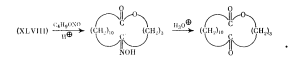
When (XLV) is irradiated with ultraviolet light in the presence of CCl₄, a dichlorovinyl ketone is formed³



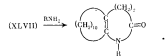
Oxopentade canolide has been synthesised from (XLV) and ethyl acrylate via the mechanism $^{\rm 369}$



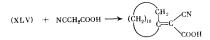
Hydrolysis of (XLVII) leads to β -(2-oxocyclododecyl)propionic acid³⁴⁴,³⁷⁰. Ultraviolet irradiation of this acid results in the formation of ω -formyl-4-tetradecenoic acid, whose reduction leads to ω -hydroxypentadecanoic acid³⁷⁰. Oxopentadecanolide has also been obtained from (XLVIII) by treatment with butyl nitrite and subsequent hydrolysis³⁷¹:



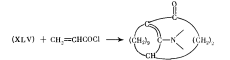
Similarly 1-acetyl-13-hydroxyimino-1-azahexadecan-2-one was obtained from 1-acetyl-2,3-decamethylene-1,4,5,6tetrahydropyridine³⁷². Treatment of the ketoester obtained from (XLVII) with alkylamines leads to the synthesis of bicyclic lactams³⁷³:



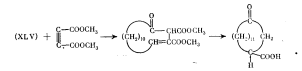
The reactions of (XLV) with a substituted cyclopropene³⁷⁴ and dibromoacetone have been investigated³⁷⁵. The morpholine residue in (XLV) undergoes an exchange reaction with cyanoacetic acid³⁷⁶:



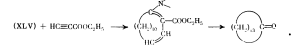
When (XLV) interacts with acryloyl chloride, a bicyclic product is formed 377 :



 α -(2-Oxocyclododecyl)acetic acid has been obtained by the reaction of (XLV) with a bromoacetate ester³⁴⁴. When pyrrolidinocyclododecene condenses with an ester of ace-tylenedicarboxylic acid³⁷⁸, ring expansion takes place with formation of a cyclotetradecenone derivative, which has been converted into 3-carboxycyclotetradecanone³⁷⁶:

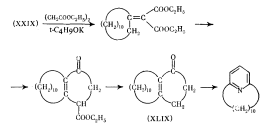


Cyclotetrade canone has been obtained by the reaction of (XLV) with ethyl propiolate³⁷⁹:

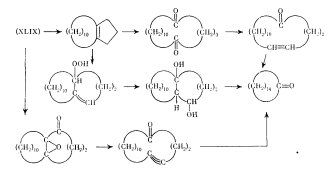


Using this reaction, it is possible to obtain the racemic muscone³⁸⁰. A series of methods of synthesis of muscone, cyclopentadecanone (exaltone), pentadecanolide (tibetolide), and methylpentadecanolide (muscolide), which are valuable perfumes, have been developed on the basis of cyclodode-canone.

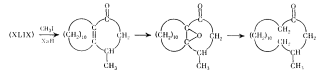
The synthesis of natural muscopyridine from cyclododecanone has been achieved by the Stobbe reaction via $(XLIX)^{381}$:

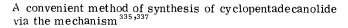


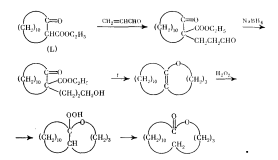
This reaction has served as the basis of a method of synthesis of cyclopentade canone $^{382-385}$:



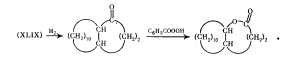
Muscone has been obtained from (XLIX) via the mechanism 383







has been developed. The methyl analogue of cyclopentadecanolide (muscolide) has been obtained similarly from (L) and methacrolein ^{328,337}. It has been reported that γ , δ -decamethylene- δ -valerolactone has a strong musk-like scent ^{386,387}. It has been synthesised via the mechanism



The products of the reaction of cyclododecanone with glycols³⁸⁸ are components of perfumes.

During the preparation of the review for the press, studies which are of interest were published. An industrial method of synthesising compound (II) (1,5,9-CDT) by a continuous method in the presence of the $(C_2H_5)_2AlCl-$ TiCl₄ catalytic system has been developed ³⁸⁹. A mechanism has been proposed for the cyclisation of butadiene to (I) in the presence of $Ni(C_5H_7O_2)_2 + (C_2H_5)_3A1.^{390}$ Partial ozonolysis of (I), with subsequent reduction of the peroxycompounds formed by dimethyl sulphide, leads to 1,12-diformyl-trans, trans-4,8-dode cadiene³⁹¹. Conditions have been found for the selective hydrogenation of (II) to (XXVI) in the presence of $RuCl_2(PPh_3)_3$ activated by $(C_2H_5)_3N$.³⁹² The reaction of (XXV) and (XXVI) with Nbromosuccinimide in the presence of water, methanol, or acetic acid proceeds with formation of 2-bromocyclododecanol, 2-bromo-1-methoxycyclododecane, and 1-acetoxy-2-bromocyclododecane respectively³⁹³. Oxidation of a mixture of (XXV) and (XXVI) by Jones reagent in the presence of mercury propionate leads to cyclododecanone It was noticed for the first time that oxidation of cyclododecane by air results in the formation of 2-3% of transepoxycyclododecane³⁹⁵. The isomerisation of epoxycyclododecane to cyclododecanone proceeds with a high yield in the presence of palladium and rhodium catalysts³⁹⁶. Whe When epoxycyclododecane is acted upon by formaldehyde in the presence of AlCl₃, 13,15-dioxabicyclo[10,3,0]pentadecane and 13,15,17-trioxabicyclo[10,5,0]heptadecane are formed ³⁹⁷. On ultraviolet irradiation, a mixture of 36% of methyl cycloundecanecarboxylate, 17% of $\mathit{cis-}$ and 25%of trans-cyclododecenones, and 9% of cis-3-cyclododecenone is formed on ultraviolet irradiation of α -diazocyclododecanone³⁹⁸; the reaction involving 2-methylcyclododecanone takes place with ring cleavage and the formation of 40% of formyl-12-dodecene³⁹⁹. A synthesis of bicyclo-[10,3,0]pentadeca-1(12)-en-13-one and its methyl analogues from cyclodode canone has been proposed 400. Formylcyclodecane has been obtained from cyclododecanone and ethoxymethylmagnesium chloride⁴⁰¹. Treatment of 1,5cyclododecadiene with N-bromosuccinimide and subsequent reduction of the product with LiAlH₄ lead to 13-oxabicyclo-[8,2,0]dodecane⁴⁰².

A method of synthesis of cyclododecanone oxime on heating cyclododecanone with ammonia in the presence of t-butyl hydroperoxide has been described in a patent⁴⁰³. A method of synthesis of 1,3- and 1,4-cyclododecanediones from a mixture of *cis*- and *trans*-3-cyclododecenones has been discovered⁴⁰⁴. [10]Paracyclophane has been synthesised from cyclododecene and methyl acetylenecarboxylate⁴⁰⁵. [9](2,4)Pyrrolophane has been synthesised in two ways: (1) by the condensation and cyclisation of 3-formylcyclododecanone with ammonia and (2) by the condensation of chlorocyclododec-2-en-1-one with di(ethoxycarbonyl)methylamine⁴⁰⁶.

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