PHOTOCHEMICAL TRANSFORMATIONS OF NATURAL PRODUCTS

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Photochemical transformations deserve the same status as any of the other standard reactions of Organic Chemistry. By judicious selection of the chromophore involved, and of the reaction conditions, high yields of defined products can be obtained. There are many examples of new photochemical reactions in the recent literature and some of these have been used to good synthetic purpose. In our view any degradative or synthetic problem in modern Organic Chemistry deserves at least consideration in photochemical terms.

The present lecture, in fact, has only a minor component of photochemistry, but the content should be of interest to Czech chemists, since the compounds discussed are the only representatives of nine-membered carbon-rings in natural products apart from caryophyllene. The presence of a nine-membered ring in the latter hydrocarbon was, of course, first proposed by Professor Sorm, Dr Herout and their co-workers nearly fifteen years ago. The lecture is concerned with the chemistry of glauconic, glaucanic and byssochlamic acids. The work described has been carried out in collaboration with Messrs J. E. Baldwin, J. L. Bloomer, L. D. S. Godinho, Professor L. M. Jackman (now of Melbourne), Drs Lydia Rodriguez-Hahn (now of the University, Mexico City) and J. K. Sutherland.

Glaucenic and glaucanic acids were first isolated by Wijkman from a mould, described as Penicillum glaucum, which has since been reclassified as Penicilliun purpureogenum. The chemistry of glauconic acid, C_{18}H_{29}O_{7}, was extensively studied at Munich by Wijkman, Kraft, Sutter and their co-workers who showed that it contained an acylable hydroxyl group and titrated as a tetracarboxylic acid. Reduction with zinc dust and acetic acid gave a dihydro-derivative to which we make further reference below. The main effort of the earlier workers was concentrated on the pyrolysis products of glauconic acid. On heating to 200° the molecule breaks down into diethylacrolein (I), C_{6}H_{12}O, identified as the corresponding acid, and glauconin, C_{11}H_{14}O_{6}. This latter compound was studied in some detail.

Glaucanin titrated as a tetracarboxylic acid, and on ozonolysis gave more than one mole of pyruvic acid as well as oxaloacetic acid. One of the most informative experiments was the reduction of glauconin with red phosphorus and hydriodic acid to a dihydro-derivative, isolated as a compound of the composition (C_{11}H_{19}O_{6} + H_{2}O). The latter was, in fact, a dicarboxylic acid, which, on conversion to the dimethyl ester with diazomethane and ozonolysis, gave (after hydrolysis of the product) the tricarboxylic acid...
acid (II). The constitution of this tricarboxylic acid was confirmed by synthesis.

When glauconin was heated with hydrochloric acid at 200° it afforded a mixture of the meso- and racemic forms of the keto-dicarboxylic acid (III) as well as the three possible racemates of the dilactone (IV). The keto-dicarboxylic acid (III) was synthesized by double alkylation of acetone-dicarboxylic ester with ethyl α-bromopropionate to give the tetra-ester (V). Hydrolysis of the latter gave the desired keto-diacid (III) which with acetyl chloride cyclized to the dilactone (IV). Mainly on the basis of this evidence the constitution (VI) was proposed for glauconin. Although the alternative (VII) has been considered, and rejected, by the earlier workers it appeared to us to explain the chemistry of glauconin better than (VI). In the event, the correctness of the formula (VII) has been conclusively established by spectral evidence and by synthesis.

The ultra-violet spectrum of glauconin in non-hydroxylic solvents showed \( \lambda_{\text{max}} \) 250 m\( \mu \) (\( \epsilon = 10,200 \)) indicative of two dialkylmaleic anhydride residues. The infra-red spectrum showed two five-membered anhydride rings. This conclusion was confirmed by inspection of the infra-red spectrum of glauconin dimethyl ester (VIII), a compound obtained by methylating glauconin with dimethyl sulphate under alkaline conditions. The N.M.R. (nuclear magnetic resonance) spectrum of glauconin in trifluoroacetic acid showed the presence of two identical unsplit methyl groups (\( \tau = 7.70 \)) attached to double bonds. The other two protons showed a single peak at \( \tau = 6.26 \) in agreement with the presence of a methylene group flanked by two ethylenic linkages.
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The constitution of glaucinin as (VII) was confirmed by synthesis according to the following method. The known tetra-ester (V) (see above) was converted by refluxing in benzene in the presence of methanesulphonic acid (azeotropic removal of water and ethanol) into a mixture of the lactones (IX) and (X). The enol-lactone was the main component of the mixture. On alkaline hydrolysis it afforded the keto-dicarboxylic acid (III). The second ester (X) gave, by β-elimination, prototropic migration and hydrolysis, glaucinin (VII) identical with authentic material. The synthesis is essentially a reversal of the hydrochloric acid induced degradation of glaucinin to the keto-dicarboxylic acid (III).

We now turn to the constitution of glauconic acid, \([\alpha]_D + 33^\circ \ (c, 1\cdot02), \lambda_{\text{max}} 223 \text{ m}_{\mu} (\epsilon = 10,500), \) inflexion near 260 m\(_\mu\), \(\nu_{\text{max}} 3510, 1830, 1770\) and 1670 cm\(^{-1}\). It can be seen that the infra-red spectrum confirms the presence of an hydroxyl group and suggests that the molecule contains one or more cyclic five-membered anhydride functions probably conjugated with ethylenic linkages. The presence of two anhydride systems was conclusively established by derivatives described in the sequel. Using the \(\Delta^1\)- and \(\Delta^2\)-tetrahydrophthalic anhydrides as models for the possible chromophores formed from cyclic anhydrides and ethylenic linkages, it was found that the ultra-violet absorption spectrum of glauconic acid was best accounted for by the presence of one \(\Delta^1\)-type chromophore \(\lambda_{\text{max}} \text{cyclohexane} 250 \text{ m}_{\mu} (\epsilon = 3,540)\) and one \(\Delta^2\)-type chromophore \(\lambda_{\text{max}} \text{cyclohexane} 223 \text{ m}_{\mu} (\epsilon = 7,600)\). The ultra-violet spectra of various derivatives (see below) have confirmed this conclusion.

As briefly noted above, reduction of glauconic acid with zinc dust in refluxing acetic acid gives\(^4\) a dihydro-derivative, \([\alpha]_D + 57^\circ \ (c, 1\cdot01) \lambda_{\text{max}} 230 \text{ m}_{\mu} (\epsilon = 8,000), \nu_{\text{max}} 1835, 1767\) and 1720 cm\(^{-1}\), which must contain an itaconyl type of anhydride chromophore. That this reduction involves more than the saturation of an ethylenic linkage is obvious from the fact that the compound is a carboxylic acid, giving with diazomethane, a monomethyl ester \([\alpha]_D + 34^\circ \ (c, 1\cdot21), \lambda_{\text{max}} 232 \text{ m}_{\mu} (\epsilon = 8,550), \nu_{\text{max}} 1840, 1780\) and 1740 cm\(^{-1}\). This methyl ester shows no hydroxyl absorption in its infra-red spectrum. The simplest relationalization of these facts is that the maleic anhydride residue has been reduced and the reduced anhydride ring opened with concomitant lactonization. On treatment with dimethyl sulphate and alkali dihydroglaucunic acid gives\(^7\) a trimethyl ester, the anhydride ring being opened and methylated. This ester shows, \(\nu_{\text{max}} 1775\) and 1745 cm\(^{-1}\), indicating the presence of a γ-lactone grouping as well as of the three ester residues. The part-formulae (XI) and (XII) will explain this transformation. The former (XI) is preferred for two reasons. First, reduction of glauconic acid acetate\(^1\), \([\alpha]_D + 35^\circ \ (c, 1\cdot20), \lambda_{\text{max}} 221 \text{ m}_{\mu} (\epsilon = 10,600), \) inflexion at about 250 m\(_\mu\), \(\nu_{\text{max}} 1835, 1770\) and 1740 cm\(^{-1}\), with zinc dust in refluxing acetic acid gave glaucanic acid, \(\text{C}_{17}\text{H}_{26}\text{O}_8, \ [\alpha]_D + 185^\circ \ (c, 1\cdot05), \lambda_{\text{max}} 220 \text{ m}_{\mu} (\epsilon = 10,700), \) inflexion at about 250 m\(_\mu\), \(\nu_{\text{max}} 1840\) and 1775 cm\(^{-1}\). As would be expected from theoretical considerations developed later, glaucanic acid, itself, is reduced to a dihydroglaucanic acid on more prolonged reduction\(^6\). Glaucanic acid must be desoxyglaucanic acid. The removal of the acetyl residue under such mild conditions is most easily understood if it is allylic to the maleic
anhydride type residue. Secondly, the N.M.R. spectrum of gluconic acid acetate shows that there is a hydrogen atom attached to the carbon atom bearing the acetoxy group and that this is split into a doublet ($\tau = 4.15$; $J = 5$ c/s). There is, therefore, only one proton on the neighbouring (z) carbon atoms. However, in dihydrogluconic acid methyl ester (see above) this same proton now appears as a triplet ($\tau = 5.06$; $J = 4$ c/s) so that it must be split by two protons. Only if we base our constitutional arguments on formula (XI) is there the requisite increase in the number of neighbouring protons on reduction. The change in the position of absorption of the proton is also in agreement with the postulated structural change. The part-formula (XI) can now be expanded to (XIII) on the basis of these arguments.

Gluconic acid can be smoothly oxidized to a ketone, C$_{18}$H$_{16}$O$_7$, m.p. 174–176°, $[\alpha]_D + 45° (c, 0.56)$, $\lambda_{max} = 220 m\mu$ ($\epsilon = 10,000$), inflexion at about 290 m\mu, $\nu_{max} 1830, 1770$ and 1695 cm$^{-1}$, with chromium trioxide in acetone. This ketone is readily reduced under the usual zinc dust-acetic acid conditions to a keto-tricarboxylic acid, C$_{17}$H$_{24}$O$_7$, m.p. 171–173° (decomp.) $[\alpha]_D - 90° (c, 1.00)$, $\epsilon_{220 m\mu} = 600$, $\nu_{max}$ 1750 and 1735 cm$^{-1}$, readily converted by diazomethane into a keto-trimethyl ester, C$_{20}$H$_{36}$O$_7$, m.p. 106–108°, $[\alpha]_D - 101° (c, 1.00)$, $\epsilon_{220 m\mu} = 500$, $\nu_{max}$ 1735 cm$^{-1}$. On heating, the tricarboxylic acid gave an anhydride, C$_{17}$H$_{22}$O$_6$, m.p. 168–170°, $[\alpha]_D - 59° (c, 1.00)$, $\epsilon_{220 m\mu} = 550$, $\nu_{max}$ 1860, 1785, 1750 and 1720 cm$^{-1}$. The absorption in the infra-red due to the ketonic carbonyl group is best resolved in this anhydride and indicates (at 1750 cm$^{-1}$) that the compound is a cyclopentanone. Furthermore, from the lack of ultraviolet absorption it must be concluded that besides reduction of the maleic anhydride type residue and decarboxylation, the itaconic anhydride type residue has also disappeared. This can only be explained by cyclization. A monocyclic substance, the ketone, must therefore have been cyclized to a bicyclic compound. Since the ketonic carbonyl was necessary for this reaction to take place, the most simple explanation was that an intramolecular Michael reaction between one of the positions $\alpha$ to the ketone group and the $\beta$-position of the itaconic residue had occurred. When the maleic anhydride residue in the ketone, depicted in (XIV), is reduced there are two $\alpha$-positions (on either side of the ketone) which could act as anion sources for the Michael reaction (see XV). However, it must be the position between the two carbonyl groups in (XV) which is involved for the following reason. During the reduction of glauconic acid ketone with zinc a second product is formed. This is a monocyclic anhydride-acid, C$_{17}$H$_{22}$O$_6$, m.p. 170–180°, $[\alpha]_D + 8° (c, 1.00)$, $\lambda_{max} = 227 m\mu$ ($\epsilon = 7,100$), $\nu_{max}$ 1830, 1770 and 1700 cm$^{-1}$, characterized as its methyl ester, C$_{18}$H$_{24}$O$_6$, m.p. 115–117°, $[\alpha]_D + 0° (c, 1.00)$, $\lambda_{max} = 225 m\mu$ ($\epsilon = 8,300$), $\nu_{max}$ 1830, 1770, 1725 and 1670 cm$^{-1}$. Obviously this compound retains the itaconic anhydride type residue intact but one carbonyl group has been lost. The maleic anhydride type residue has also been reduced in the normal way. This anhydride–acid does not cyclize to the bicyclic derivative under the conditions of the zinc dust–acetic acid reduction. One must conclude, therefore, that cyclization precedes the loss of the carboxyl group and that the latter is mandatory for cyclization. This can only be understood if the
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1:3-dicarbonyl system of (XV) (or its hydrated equivalent) is required for the cyclization. Indeed, decarboxylation may be synchronous with cyclization. In any case the "anion" for the cyclization must be on the carbon atom between the 1:3-dicarbonyl system of (XV). We can now join the maleic anhydride type residue to its itaconic analogue in such a manner that the postulated cyclization will place the ketonic group in a five-membered ring. The part formula (XIII) can therefore be expanded to (XVI), certain structural conclusions justified in the sequel being also anticipated.

The integrated N.M.R. spectrum of glauconic acid acetate shows the presence of one olefinic proton ($\tau = 3.11; J = 13 \text{ c/s}$) split as a doublet. The doublet is retained at the same position in the ketone from glauconic acid. This excludes any formula with the hydroxyl group allylic to the itaconic anhydride type residue. If the latter had been correct, the formation of the ketone would have removed the proton splitting the vinylic proton. The possibility that the proton which is causing the splitting of the vinylic proton is $\alpha$- to the carbonyl of the itaconic anhydride type residue, i.e. that coupling is taking place across the double bond, is excluded by the magnitude (13 c/s) of the coupling constant. This conclusion is already allowed for in part formula (XVI).

The nature of the two carbon bridge between the vinylic CH and CHOH groups of (XVI) was established by the isolation of meso-diethylsuccinic acid from oxidation experiments (see below) and supported by the isolation of diethyl-acrolein from the pyrolysis of glauconic acid. Part formula
(XVI) can therefore be expanded to (XVII). The latter contains 19 carbon atoms and, therefore, one atom in (XVII), indicated by a C, must, in fact, be allowed for twice over. The final structural detail was settled quickly by the following considerations. Glauconic acid and its derivatives, where appropriate, show one quaternary C-methyl group in their N.M.R. spectra. Formula (XVII) must, therefore, be modified to (XVIII; R = H) as the unique constitution of glauconic acid. The methylene group in this formula can be identified in the derived glauconic acid ketone (XIX) as an AB quartet ($\tau = 6\cdot677, 6\cdot96$; J = 15 c/s). It must, therefore, be placed between two fully substituted carbon atoms as already allowed for in (XIX) and related formulæ.

We may now consider the chemistry of glauconic acid in the light of the constitution (XVIII; R = H) now established. It is obvious that this formula does not contain the glauconin carbon skeleton and that the formation of this substance, in fact, gives a misleading clue to the constitutional problem. It seems that glauconic acid must first undergo a Cope$^{12, 13}$ rearrangement (XVIII; R = H; see arrows) to (XX; R = H) which is then converted to (XXI) by a reversed aldol reaction. The degradation of (XX) to diethylacrolein (I) and glauconin (VII) then follows by a conventional reversed Michael reaction.

![Chemical structures](image)

The pyrolysis of glauconic acid acetate (XVIII; R = Ac) provides support for these arguments. There is formed an isomeric compound, isogluconic acid acetate, m.p. 257–258°, [a]$_D$ = 5° (c, 1·36), $\lambda_{max}$ 220 m\(\lambda\), $\epsilon$ = 10,500; $\nu_{max}$. 1835, 1776 and 1742 cm$^{-1}$, which contains one ethylenic linkage, since it furnishes on hydrogenation over 10 per cent palladized charcoal a dihydroisogluconic acid acetate, m.p. 195–202°, [a]$_D$ = 104° (c, 0·52), $\epsilon_{220 m\lambda}$ = 300, $\nu_{max}$. 1850, 1780 and 1715 cm$^{-1}$. This derivative is saturated and therefore has two carbocyclic rings. Isogluconic acid acetate has two vinyl protons ($\tau = 3\cdot26$ and 3·95) each split

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as a doublet with a coupling constant of about 1 c/s. This evidence, supported by that of the ultra-violet absorption spectrum, indicates the presence of an itaconic anhydride residue with the ethylenic linkage terminal (as =-CH=). This was confirmed by the isolation of a good yield of formaldehyde on ozonolysis. If glauconic acid acetate rearranges on pyrolysis to (XX; \(R = \text{Ac}\)) then a further rearrangement can be envisaged (XX; \(R = \text{Ac}\); see arrows) which would give the formula (XXII; \(R = \text{Ac}\)) for isoglauconic acid acetate. This formula would agree with the fact (N.M.R.) that the compound has no C-methyl groups (apart from those in the two ethyl groups).

The zinc-acetic acid reduction of glauconic acid to dihydroglauconic acid has already been referred to above. This reaction can be understood if one considers that the reaction takes the course shown below.

![Diagram](https://example.com/diagram.png)

The donation of electrons from the zinc metal affords an anion (in the \(\beta\)-position) which is stabilized since it is \(\alpha\)- to a carbonyl group. A similar reduction of an itaconic anhydride type unit through carbonyl oxygen is not probable since the carbanion formed would not be stabilized in the same way. Isomerization of the initially formed succinic anhydride by interaction with the allylically placed hydroxyl group would then afford the \(\gamma\)-lactone-acid, dihydroglauconic acid (XXIII). In the case of glauconic acid, acetate competition with anionic \(\beta\)-elimination of acetoxy can be envisaged to give glauconic acid (XXIV). The N.M.R. spectrum of glauconic acid is in full accord with this formula.

The reduction of glauconic acid ketone with zinc dust and acetic acid has already been discussed at some length above. The saturated bicyclic ketone thereby produced can now be formulated as (XXV), whilst its companion unsaturated ketone can be represented as (XXVI). Further support for formula (XXV) has been secured by dehydrogenation over palladized charcoal which afforded an aromatic ketone, \([\alpha]_D \pm 0^\circ\), characterized as an \(\alpha\)-hydrindanone by its ultra-violet (\(\lambda_{\text{max}}\) 250 m\(\mu\) (\(\epsilon = 8,100\)), 274 m\(\mu\) (\(\epsilon = 2,500\)) and 294 m\(\mu\) (\(\epsilon = 1,800\))) and infra-red (\(\nu_{\text{max}}\) 1700 cm\(^{-1}\)) spectra. This compound should have formula (XXVI), which constitution has been confirmed by synthesis.

Pyrolysis of dihydroglauconic acid chloride at 200\(^\circ\) for 30 min gave an isodihydroglauconic acid, \(C_{18}H_{22}O_7\), m.p. 227–230\(^\circ\), \([\alpha]_D + 5^\circ\) (\(\epsilon = 1.00\)), \(\epsilon_{220m\mu} = 500\), \(\nu_{\text{max}}\) 1860, 1785 and 1730 cm\(^{-1}\), characterized as the monomethyl ester, \(C_{19}H_{24}O_7\), m.p. 165–167\(^\circ\), \([\alpha]_D + 12^\circ\) (\(\epsilon = 0.90\)), \(\epsilon_{220m\mu} = 800\), \(\nu_{\text{max}}\) 1860, 1785 and 1730 cm\(^{-1}\). This compound, which is saturated and therefore bicarbocyclic, is formulated as (XXVIII), being
formed by a Michael type reaction between the $\alpha$-position of the acid chloride and the $\beta$-position of the itaconic anhydride type residue.

Reference to several of the reactions of glauconic acid ketone (XIX) has already been made in the text above. This ketone is an extremely sensitive compound which is attacked by many reagents. For example, if the ketone is refluxed with cyclohexylamine carbonate in benzene it affords an amide, $C_{23}H_{31}O_5N$, m.p. 178–179°, $[\alpha]_D = 26^\circ (c, 1.06)$, $\lambda_{\text{max}}. 218 \text{ m}\mu (\epsilon = 12,700)$, $\nu_{\text{max}}. 1830, 1775, 1690$ and 1650 cm$^{-1}$, whose properties are in accord with the constitution (XXIX). The N.M.R. spectrum of this amide shows two vinyl protons, one as a doublet ($\tau = 3.09; J = 10 \text{ c/s}$) and another as a singlet ($\tau = 3.44$). On the other hand, if glauconic acid ketone (XIX) is reacted under the same conditions with pure cyclohexylamine an amide-imide, $C_{27}H_{38}O_5N_2$, m.p. 195–197°, $[\alpha]_D = 28^\circ (c, 0.99)$, $\lambda_{\text{max}}. 226 \text{ m}\mu (\epsilon = 15,600)$, $\nu_{\text{max}}. 1755, 1700$ and 1630 cm$^{-1}$, is formed. The constitution of this compound is still under investigation.

Oxidation of glauconic acid ketone with chromic acid in acetic acid gave meso-diethylsuccinic acid, the identity of which was carefully confirmed. Oxidation of glauconic acid itself under the same conditions afforded the same diethylsuccinic acid and a crystalline dicarboxylic acid, $C_{18}H_{26}O_7$, m.p. 202–204°, $[\alpha]_D + 61^\circ (c, 1.20)$, $\lambda_{\text{max}}. 224 \text{ m}\mu (\epsilon = 7,450)$, $\nu_{\text{max}}. 1845, 1750, 1730$ and 1690 cm$^{-1}$. This compound clearly retains the
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itaconic anhydride type residue. It is formulated as (XXX; R = H). Treatment with diazomethane furnished the dimethyl ester (XXX; R = Me), C_{17}H_{32}O_{2}, m.p. 91–92°, [,]D + 55° (c, 1.30), λ_max. 223 μm. (ε = 9,500), ν_max. 1830, 1770 and 1730 cm⁻¹. The N.M.R. spectrum of this ester showed the usual vinyl proton doublet at τ = 3.28 (J = 11 c/s), the quaternary methyl at τ = 8.37 and methylene absorption (a singlet at 6.77). Further chromic acid oxidation of the diacid (XXX; R = H) gave meso-diethylsuccinic acid in trace amounts (paper chromatography).

We now turn to consideration of the stereochemistry of glauconic acid. In all the formulae used so far the vinylic hydrogen of glauconic acid (XVIII; R = H) has been written cis with respect to the anhydride residue. This assignment is made¹⁴a because the average position of the vinylic proton in the N.M.R. spectra is at τ = 3.10. The vinylic proton in Δ²-tetrahydrophthalic anhydride absorbs at τ = 2.95. The best compound for comparison is glauconic acid (XXIV) whose vinylic proton absorbs at τ = 3.00. In itaconic anhydride the two vinylic protons (one cis, one trans with respect to the anhydride) differ in absorption by 0.43 p.p.m. The close agreement between the position of absorption of the vinylic proton of glauconic acid (XXIV) and the model Δ²-tetrahydrophthalic anhydride provides then good support for the assigned cis-configuration.

The relatively large coupling constant for the splitting of the vinylic proton of glauconic acid and its derivatives by the adjacent allylic hydrogen has already been mentioned (see above). This implies⁹ that the dihedral angle between the two C—H bonds involved must be about 180°. Examination of molecular models suggests that this condition can only be satisfied if the angular methyl group at C-15 and the ethyl group at C-12 are trans to each other. The degradation of the molecule to meso-diethylsuccinic acid shows that, provided epimerization has not occurred during the reaction sequence, the two ethyl groups at C-12 and C-3 must be cis to each other in the nine-membered ring. There remains only the configuration of the hydroxyl group for consideration. Definitive chemical evidence on this point has not yet been obtained. These tentative conclusions as to the stereochemistry of glauconic acid are summarized (without commitment as to absolute configuration) in the formulae (XXXI; R = OH) for glauconic acid and (XXXI; R = H) for glauconic acid. We have prepared the m-iodobenzoate of glauconic acid, C_{25}H_{32}IO_{5}, m.p. 197–198°, [,]D + 89° (c, 0.64), ν_max. 1840, 1770 and 1725 cm⁻¹. This substance has been examined by the X-ray method by Professor J. M. Robertson, Dr G. A. Sim, and their co-workers¹⁵ in order to define with certainty the stereochemistry of glauconic acid. The tentative chemical conclusions on the stereochemistry have been confirmed. The hydroxyl group of glauconic acid is in the β-configuration (see XXXI, R = OH).

Our investigations on the absolute configuration of glauconic acid are not yet complete. We can, however, outline briefly the method on which we are currently working. The dicarboxylic acid (XXX, R = H) is being degraded to the bisnoriodide by the use of lead tetra-acetate and iodine under illumination with visible light. This is a procedure which we have developed¹⁶ for the decarboxylation of primary and secondary carboxylic acids in high yield. Reduction of the bis-iodide by zinc and then ozonolysis should afford
optically active α-ethyl valeric acid. From the rotation of this compound it
should be possible to deduce the absolute configuration of gluconic acid.

It is now opportune to discuss the biogenesis of gluconic and glaucanic
acids. In common with many other fungal metabolites we have shown in
preliminary experiments that they are derived from acetate\(^{17}\) (or the equiva-
lent malonate\(^{18}\)). However, there is an aspect of the constitutions of these
compounds which we consider makes them of unusual biogenetic interest.
In principle, the constitution of gluconic acid (XXXI, \(R = \text{OH}\)) may be
derived from two units having identical carbon skeletons (XXXII). This
we have already implied in the numbering system adopted. The oxidation
level of gluconic and glaucanic acids is such that if, in principle, one took
the dienoid unit (XXXII), an anion from one molecule could serve as an
agent for the construction of the whole carbon skeleton of glaucanic acid with
the substituent ethylene linkages in the correct positions (see XXXIII).
There are, of course, many equivalent representations. If this hypothesis be
granted then the hydroxyl group of gluconic acid would be introduced at a
later stage in the biogenesis, possibly by biochemical hydroxylation of
gluconic acid.

The carbon skeleton of (XXXII) is readily derivable from an inter-
mediate citric acid and it is possible that acids of this kind are intermediates
in the biogenesis of such compounds as lichesterenic acid (XXXIV)\(^{19}\), mine-
luteic acid (XXXV)\(^{20}\) and many other analogous fungal products. Alterna-
tively, the dienoid unit (XXXII) may be derived from condensation of a
C–6 acetate derived chain with pyruvic acid. We are at present engaged on
a study of the biogenesis of gluconic acid from \(C_2\) and higher units to test
the hypotheses here enunciated.

The literature describes the isolation and characterization of an isomer of
gluconic acid, byssochlamic acid, which also contains two anhydride rings.\(^{21}\)
Byssochlamic acid is obtained from Byssochlamys fulva Ollier and Smith.
From the outset we suspected that glaucanic and byssochlamic acids
were related biogenetically. Byssochlamic acid showed \(\lambda_{\text{cyclohexane}} 244 \text{ m}\mu\)
(\(\epsilon = 9,100\)), \(\nu_{\text{max}}\). 1845 and 1770 \text{ cm}^{-1} and showed no vinyl protons in its
N.M.R. spectrum. This evidence indicated to us that the molecule contained
two maleic anhydride residues and must, therefore, be monocarbocyclic.
On Kuhn–Roth oxidation byssochlamic acid gave two moles of volatile
acid. The N.M.R. spectrum showed that neither of the C-methyl groups
which must, therefore, be present was quaternary and made it probable
that two ethyl or higher alkyl groups were contained in the molecule. Now
it had been demonstrated in unpublished work by Cook, Loudon and
Paton\(^{22}\) that on heating to 200° byssochlamic acid was converted into an
isomer, isobyssochlamic acid, \(C_{18}H_{29}O_7\), m.p. 153°, \([\alpha]_D + 27^\circ (c, 1-00)
\lambda_{\text{cyclohexane}} 236 \text{ m}\mu\) (\(\epsilon = 6,300\)), \(\nu_{\text{max}}\). 1855, 1830, 1780 and 1770 \text{ cm}^{-1},
characterized (by dissolution in alkali and acidification) as the hydrate, a
dicarboxylic acid anhydride, \(C_{18}H_{22}O_7\), m.p. 173–183° (decomp.),
\([\alpha]_D + 24^\circ (c, 1-30), \lambda_{\text{max}}\). 234 \text{ m}\mu\) (\(\epsilon = 3,300\)), \(\nu_{\text{max}}\). 1840, 1780, 1700
and 1690 \text{ cm}^{-1}. At the melting point this hydrate gave back isobyssochlamic
acid. Hydrogenation of isobyssochlamic acid in acetic acid over 10 per cent
palladized charcoal gives\(^{22}\) the saturated dihydroisobyssochlamic acid,
\(C_{18}H_{33}O_6\), m.p. 130°, \([\alpha]_D -16^\circ (c, 0-89), \epsilon_{220\text{mu}} = 245, \nu_{\text{max}}\). 1840 and

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1775 cm⁻¹, which also gives a stable hydrate, C₁₈H₂₄O₇, m.p. 190–200° (decomp.). From this evidence it is clear that the conversion of byssochlamic to isobyssochlamic acid is a cyclization of a mono- to a bi- carbocyclic system. Furthermore, dehydrogenation of byssochlamic acid over platinum at 300° affords a compound, C₁₈H₁₈O₃, m.p. 91°, [α]D ± 0°. We have shown that this compound has λ_cyclohexane 224 and 232 μ (ε = 26,000 and 18,000 respectively), ν_max. 1835 and 1770 cm⁻¹, and that it contains two C-methyl groups by Kuhn–Roth determination. The N.M.R. spectrum showed that both these methyl groups were present as ethyl or higher alkyl groups and revealed the presence of one aromatic proton at τ = 2-72. These facts establish that the compound, C₁₈H₁₈O₃, must be a substituted phthalic anhydride, the position of the aromatic proton in the N.M.R. spectrum suggesting that it is separated from the anhydride residue by one alkyl group. Dehydrogenation of isobyssochlamic acid gave the same substituted phthalic anhydride in improved yield. Assuming that no extra rearrangement is involved in this dehydrogenation, then isobyssochlamic acid must contain a six-membered ring bearing an itaconic anhydride type residue. A second carboxylic ring must be present and, since this occupies two positions in ortho-relationship on the benzene ring of the C₁₆H₁₈O₃ compound, we thought that it was probably a five-membered ring. The isolation of benzenepentacarboxylic acid from nitric acid oxidation of the C₁₆H₁₈O₃ confirms that five substituents are attached to this benzene ring. There must be one alkyl group attached to the benzene ring of the C₁₆H₁₈O₃ compound (see above) and since this furnishes one mole of volatile acid on oxidation it must be at least an n-propyl group. Assuming that any tetralin-type compound would have dehydrogenated further, we concluded that the C₁₆H₁₈O₃ compound was a hydridane with an anhydride residue and an n-propyl group attached to the benzene ring. By exclusion there must be one ethyl group attached to the cyclopentane ring and, therefore, all the properties of the C₁₆H₁₈O₃ compound can be summarized in the expression (XXXVI). The mode of formation of isobyssochlamic acid and its relationship to (XXXVI) provides good evidence that byssochlamic acid itself contains a nine-membered ring.

If one now makes the hypothesis that byssochlamic acid is formed from the same biogenetic units, (XXXII) or equivalent, by the same type of mechanism, (XXXIII) or equivalent, as operates in the biogenesis of glaucanic acid, then one can (as first pointed out by Dr J. K. Sutherland) advance a plausible formula. Attaching the units to each other as in scheme (XXXVII) (or equivalent operation) affords (XXXVIII) as the constitution of byssochlamic acid. The C₁₈H₁₈O₃ compound, if formed without complication, would then be (XXXIX) and isobyssochlamic acid, which has no vinyl hydrogen in its N.M.R. spectrum and whose ultra-violet spectrum indicates a fully substituted itaconic anhydride type residue, would be (XL, R = H), or equivalent formulation. Bromination of isobyssochlamic acid with N-bromosuccinimide gave a monobromo-derivative, m.p. 180–185°, [α]D = 31° (c, 1-00), λ_cyclohexane 240 μ (ε = 8,600), ν_max. 1830 and 1780 cm⁻¹. The N.M.R. spectrum of this bromo-compound showed a triplet at τ = 4-14 indicative of one hydrogen attached to carbon bearing bromine, the carbon atom being flanked by a
methylene group. The part structure (\(-\text{CH}_2\text{CHBr}\)) can be accommodated by the expression (XL, \(R = \text{Br}\)).

The major difficulty in accepting formula (XL) for isobysochlamic acid is that its relationship to bysochlamic acid is not, from the mechanistic point of view, very clear. Alternative formulae such as (XLI) could explain all the facts about isobysochlamic acid except that its dehydrogenation to (XXXIX) would involve a further rearrangement. Because of these difficulties we had not reached a firm conclusion as to the structure of bysochlamic acid at a time when Professor J. M. Robertson, F.R.S., Dr G. A. Sim, and their co-workers\(^{85}\) had been able to complete an X-ray determination of the structure of bysochlamic acid bis-\(\rho\)-bromophenylhydrazide. This compound has the constitution and configuration, apart from absolute configuration, depicted in (XLII) and thus proves the correctness of formula (XXXVIII). The bis-\(\rho\)-bromophenylhydrazide, m.p. 164-166°, \([\alpha]_D = 97° (c, 0\cdot30), \lambda_{\text{max}} 236 \, \mu\text{m} (\epsilon = 41,000), 292 \, \mu\text{m} (\epsilon = 3,100), \nu_{\text{max}} 1780 \text{ and } 1724 \, \text{cm}^{-1}, \text{was prepared by treating bysochlamic acid with } \rho\text{-bromophenylhydrazine in chloroform solution.}

Further aspects of the chemistry of bysochlamic acid can now be rationalized in terms of formula (XXXVIII). As with glauconic acid, reduction of bysochlamic acid with zinc dust and acetic acid gave interesting results. There was formed, after the appropriate working up procedure, a saturated "dihydrobysochlamic acid", \(C_{18}H_{22}O_6\), m.p. 120°, \([\alpha]_D = -54° (c, 1\cdot00), \epsilon_{\text{enol}} = 350, \nu_{\text{max}} 1840 \text{ and } 1775 \, \text{cm}^{-1}, \text{further characterized as the hydrate}\(^{20}\), \(C_{18}H_{24}O_7\), m.p. 241°, \([\alpha]_D + 15° (c, 0\cdot90 \text{ in acetone}), \epsilon_{\text{enol}} = 450, \nu_{\text{max}} 1830, 1780 \text{ and } 1700 \, \text{cm}^{-1}, \text{which was reconverted into its progenitor on warming with acetyl chloride. We consider that reduction of one of the anhydride systems of bysochlamic acid gives an anion which adds to the second anhydride system as illustrated in (XLIII) \(\rightarrow\) (XLIV). Accepting only formulae with fused five- and six-membered rings, three formulae (XLV, XLVI and XLVII) can be

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written for "dihydrobyschlamic acid". We are unable to distinguish between them at the present time.

The ethylenic linkages of bysschlamic acid are, not surprisingly, very resistant to attack by electrophilic reagents. We decided, therefore, to attempt to modify the maleic anhydride systems by a novel Hofmann degradation. One of the anhydride rings was converted to N-hydroxy-imide which was treated with toluene-\(\rho\)-sulphonyl chloride and then with 1\(\pi\) aqueous sodium hydroxide at 95°. This gave two ketones, C\(_{16}\)H\(_{29}\)O\(_4\), (A) m.p. 110°, [\(\alpha\)]\(_D\) -79° (\(\epsilon\), 0-90), \(\lambda\)\(_{\text{cyclohexane}}\) 252 m\(\mu\) (\(\epsilon\) = 4,600), \(\nu\)\(_{\text{max.}}\) 1860, 1760 and 1695 cm\(^{-1}\) and (B) m.p. 155°, [\(\alpha\)]\(_D\) +6° (\(\epsilon\), 0-95), \(\lambda\)\(_{\text{cyclohexane}}\) 255 m\(\mu\) (\(\epsilon\) = 5,500), \(\nu\)\(_{\text{max.}}\) 1845, 1760 and 1686 cm\(^{-1}\).

These compounds must be formed according to the sequence (XLVIII) and, therefore, have two out of the four formulae (XLIX), (L), (LI) and (LII).
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We are currently degrading these ketones further to establish their constitutions and to obtain evidence as to the absolute configuration of byssochlamic acid.

From consideration of the constitution of byssochlamic acid we expected that it would undergo photochemical cyclization on irradiation with ultra-violet light. A photoisomer was indeed formed in good yield in tetrahydrofuran. This compound, m.p. 185–210°, \([\alpha]_D + 6^\circ\) was saturated and analysed correctly for the expected isomer (LIII). Its molecular weight (332) was confirmed by mass spectrometric determination. The only evidence against the constitution (LIII) is the stability of the compound to heat (310° for 3 hours). However, the model anhydride (LIV) was smoothly dimerized to (LV) under the same experimental conditions and this compound was likewise extremely resistant to pyrolysis.

Unless stated otherwise \([\alpha]_D\) were taken in CHCl₃, ultra-violet absorption spectra in ethanol and infra-red spectra as Nujol mulls. All m.p.s were determined on the Kofler block. N.M.R. spectra were determined in CDCl₃ and \(\tau\) values given refer to the centre of multiple absorption where present. We thank Dr J. W. Lown and Mr R. G. Foster for some of these determinations.

We express our deep appreciation to Professor H. Raistrick, F.R.S., for his constant encouragement and for supplies of glauconic and byssochlamic acids. We thank Professor J. H. Birkinshaw, Mr G. Smith and Dr C. E. Stickings for very helpful advice on the culture of Penicillium purpurogenum and for facilities generously placed at our disposal. Drs J. W. Cook, F.R.S. and J. D. Loudon kindly permitted us to see, and to quote from, their unpublished experiments. Dr E. R. S. Winter of Messrs J. and E. Sturge (Birmingham) is thanked for the supply of glauconic acid at

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Pecher made constructive contributions to attempted syntheses of glauconin and we
thank them cordially. Dr R. I. Reed (Glasgow) kindly determined the M.W.s of
glauconic acid acetate and of several other compounds described in this communication.

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