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#### THE UNIVERSITY OF ALBERTA

METABOLITES OF CYATHUS AFRICANUS

AND A RELATED FUNGUS

bу

C DIRK MARINUS JOHANNES VAN SCHIE

#### A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH
IN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE
OF MASTER OF SCIENCE

DEPARTMENT OF CHEMISTRY

EDMONTON, ALBERTA
FALL, 1977

#### THE UNIVERSITY OF ALBERTA

#### FACULTY OF CHADUATE STUDIES AND RESEARCH

The undersigned of this that they have read, and recommend to the Faculty of Graduate Studies and Research, for acceptance, thesis entitled METABOLITES OF CYATHUS AFRICANUS AND A PELATED FUNGUS

submitted by DIRK MARINUS JOHANNES VAN SCHIE in partial fulfilment of the requirements for the degree of Master of Science

W.A. Ayer, Supervisor

Date 14 October 1977.

#### ABSTRACT

Cyathus africanus were separated by thin-layer chromatography. Two of these were shown to be identical with cyathin A (1) and allocyathin B (2) respectively, both of which were first isolated from Cyathus helenae.

<u>l</u>

The third metabolite, cyafrin A, was shown to possess structure 19 by spectroscopic analysis of the parent compound and its triacetyl and acetonide derivatives. Preliminary work towards the correlation of cyafrin A with cyathin A was undertaken.

The structure of the fourth metabolite, cyafrin B (21), was established by spectroscopic analysis of

19 . 21

the parent compound and its diacetyl and methyl ketal derivatives. Many other metabolites were shown to be present but were not isolated.

A preliminary investigation into the metabolites of another bird's nest fungus, Sphaerobulos stellatus, was undertaken.

V

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#### I. INTRODUCTION

In recent years a large number of bird's nest l fungi have been described, most of which have been classified as belonging to the class <u>Basidiomycetes</u>, subclass <u>Homobasidiomycetes</u>, order <u>Gasteromycetes</u>, family <u>Nidularia-ceae</u>, genus <u>Cyathus</u>.

Wilkins reported that mycelium of one of these fungi, Cyathus striatus, showed bacteriostatic properties. This report was confirmed later by Broadbent. In 1967 Olchowecki found that the mycelium of Cyathus helenae also inhibited growth of a number of bacteria. Further studies indicated that mycelium of other species, including Cyathus limbatus and Cyathus poeppigii, also exhibited bacteriostasis.

Of these fungi C. helenae has been studied most extensively. Johri found that an ethyl acetate extract of the liquid medium used for cultivation of this fungus retained its antibacterial activity. The residue from evaporation of this extract was named "cyathin". Johri also carried out studies on the effect of the composition of the nutrient medium and other factors on the growth of C. helenae and the propertion of cyathin.

Taube first separated several components of crude cyathin. He identified the phenol 2,4,5-trihydroxy-

benzaldehyde and found a large number of C compounds all 20 of which appeared to have the same basic skeleton. These compounds were named cyathins, with a letter and a number added to the name of each to indicate the number of hydrogen and oxygen atoms in the molecule, respectively. Thus compounds of the "A" series contain 30 hydrogen atoms, of the "B" series 28 and of the "C" series 26. The number in each name corresponds to the number of oxygen atoms in the molecule. If a second compound was found with the same molecular formula it was distinguished from the first by the prefix "allo-", and a third one by "neoallo-".

Taube succeeded in obtaining in pure, crystalline form two of these, cyathin A (C H O)(1) and allocyathin B (C H O)(2), and determined their structures 3 20 28 3 8,9 by spectroscopic and chemical methods. The absolute stereochemistry has been confirmed by X-ray diffraction studies of cyathin A.

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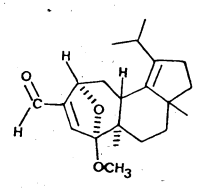
Carstens carried on the investigation of

crude cyathin. He separated a mixture of cyathin B (C H 0) (3) and cyathin C (C H 0) (4) from the re- 20 28 3 3 20 26 3 maining components of the crude cyathin. He was able to

3

4

separate them from one another only in the form of their methyl ketal derivatives ( $\underline{5}$  and  $\underline{6}$ , respectively). The struc-



5

tures of cyathin B (3) and cyathin C (4) were determined 3 from spectroscopic evidence on the methyl ketals (5 and 6 respectively) and confirmed by chemical correlation. The methyl ketal of cyathin B was converted to the methyl ketal of cyathin A and the methyl ketal of cyathin C was converted to the methyl ketal of cyathin C and the methyl ketal of allocyathin B. Later Mercer developed a direct method of separating cyathin B from 12 cyathin C .

Recent work by Mercer resulted in the isolation of neoallocyathin A (C H O) (7) via its 0,0-iso-4 20 30 4 propylidene derivative 8.

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<u>7</u>

Two other bird's nest fung; that were subsequently investigated, Cyathus bulleri and Cyathus interlated, were found not to produce any compounds of similar structure to the cyathins.

In 1966 Dr. D. Hocking brought with him from Tanzania two specimens of a bird's nest fungus which he had found growing on dead twigs on Mount Kilimanjaro at an altitude of about 8000 ft. Brodie determined this to be a previously unrecognized species of the genus Cyathus and named it Cyathus africanus.

Johri compared chromatograms of crude cyathin from <u>C</u>. helenae with those of similarly prepared extracts of <u>C</u>. striatus and of the recently discovered <u>Cyathus</u> africanus Brodie, and found them to be very similar. although not identical.

It was the similarity between chromatograms of extracts of <u>C</u>. <u>africanus</u> and <u>C</u>. <u>helenae</u> that prompted the investigations presented in this thesis. To distinguish the metabolites of <u>C</u>. <u>helenae</u> and <u>C</u>. <u>africanus</u>, the crude ethyl acetate extract from <u>C</u>. <u>africanus</u> will be called "cyafrin".

The purpose of this work was thus to determine whether extracts of <u>C</u>. <u>africanus</u> also contain compounds similar to the ones found in <u>C</u>. <u>helenae</u>, and to elucidate the structure of any new compounds isolated.

Another related fungus, Sphaerobolus stellatus, also had been found to exhibit some antibacterial activity. For this reason the metabolites of this fungus were also investigated.

#### II. GENERAL EXPERIMENTAL

## Growth of fungi and preparation of crude extracts

The procedures used for growing both  $\underline{C}$ .  $\underline{afri}$ - $\underline{canus}$  (strain 66120) and  $\underline{S}$ .  $\underline{stellatus}$  are based on those developed for  $\underline{C}$ .  $\underline{helenae}$  by  $\underline{Johri}^{\dagger}$ .

Agar slant tubes of the fungus strain 66120 were obtained from the collection of H. J. Brodie, and were maintained on Brodie solid (agar) medium (for the composition of the Brodie medium used see detailed experimental). These cultures may be stored for long periods on solid medium in slant tubes or on Petri dishes at 5 C.

The growth of mycelium in liquid medium is initiated by transferring some of the agar-grown mycelium to a 500 ml Erlenmeyer flask containing 250 ml of sterilized Brodie medium. The culture is allowed to grow at room temperature for about four weeks, then the liquid culture is homogenized in a sterilized Waring blender and used to inoculate (by means of a sterile pipette) a number of 2 l Fernbach flasks each containing 500 ml of sterile medium. After about four weeks of growth the liquid is decanted from the flasks. The remaining mycelium may be reflooded with fresh sterilized medium which will yield after a further four weeks a second yield of metabolites. The

quantity of mycelium in the flask has usually increased greatly after the second growth so that repeating the "reflooding" technique is impractical. A new growth is then initiated.

Crude extracts are prepared by partitioning twice the combined liquid media with half the volume of ethyl acetate. The organic layer is dried with anhydrous sodium sulphate, filtered, and the solvent is removed by means of a rotary evaporator.

#### Solvents and adsorbants

Because of the small scale on which separations and reactions had to be carried out during this work even small amounts of involatile impurities in solvents could not be tolerated. Therefore all solvents were distilled before use except anhydrous ethyl ether (Mallinckrodt), which was found to be of sufficient purity to be used directly from the container.

e only adsorbent used was Silica gel G

(E. Merck dsorbent was used directly from the bottle.

## Analytical this comatography (tlc)

1. Preparation of Lances

Or ym a scop slides were used, of either

25 x 75 mm or 50 x 75 mm. These were coated with the adsorbent layer with the aid of templates, using a Desaga
16
spreader, according to a method described by Stahl . Routinely a slurry was made up of 50 g silica gel G, approx.

1 g "electronic phosphor" (ZnSiO , General Electric) and
100 ml H O. The spreader was set to give a thickness of
2 adsorbent layer of 0.2 - 0.3 mm. The coated slides were
allowed to dry at room temperature for several hours and then activated by heating overnight at 110 - 120 C.

2. Application of samples and developing of the plates.

Samples were applied as solutions in a solvent as non-polar as possible, using open end melting point tubes which had been drawn out in a flame to give a very fine capillary point. Up to three spots could be applied to a small microscope slide and up to six to a large one.

Plates were developed inside a 250 ml beaker covered with a petri dish. Approximately 10 ml of the appropriate solvent was needed. Development time depended on the solvent system, but was approximately five minutes.

In certain cases the "multiple elution" technique was employed, whereby a plate, after having been developed, was allowed to dry and then returned to the developing tank to be eluted a second time. This process was
repeated as often as needed.

## 3. Detection of components.

Components could be detected either by viewing the plates under uv light (254 nm) or by spraying the plates with a 30% h SO solution followed by heating 2 4 at 110 C for a few minutes. In the first case spots would show up dark against a bright green fluorescent background, in the second case the spots would be charred to a purple or yellow-brown colour. Some indication as to the relative amounts of material in each spot could be obtained from its size and the intensity of charring.

4. Recording of R values and solvent systems used. f

R values are given as the ratio of the disf
tance moved by the component over that moved by the solvent
front. These values are dependent on the solvent system used.
This is indicated (by its code letter in parentheses) along
with the value. See below for these solvent systems.

In cases where "multiple elution" was applied the R was recorded after the final elution. The solf vent system was allowed to progress to the same point on each elution.

## Solvent systems used were:

A: Benzene: Acetone: Acetic Acid 75: 25:1

B: Chloroform : Methanol 97 : 3

C: Chloroform : Methanol 19 : 1

D: Acetone : Skellysolve B 7

#### Preparative thin-layer chromatography

1. Preparation of plates.

For preparative work, plates of size 100 x 20, 20 x 20, 10 x 20 and 5 x 20 cm were used. These were coated in a similar manner to the analytical plates, using a slurry of the same composition, with the spreader set to give a layer of  $_{\Omega}$ 0.7 - 0.8 mm thickness.

2. Application of samples and developing of plates.

For small samples the application was done by hand, using a small disposable pipette, the tip of which had been drawn out over a flame to a fine capillary, and which was fitted with a rubber bulb over the other end. Solutions were drawn up into the pipette with the rubber bulb and applied to the plate by gently squeezing the bulb while moving the pipette slowly along a line parallel to and about 2 cm from the bottom edge of the plate. For large plates the application was done with a mechanical applicator equipped was a similar capillary.

In general, it was found that 5 mg of material per cm width of plate could be separated without difficulty.

The small plates were developed in glass tanks of about  $25 \times 25 \times 5$  cm, which were closed at the top with a glass plate. The large plates were developed in a metal tank (Shandon) of  $110 \times 25 \times 15$  cm. Developing time was approximately 50 minutes.

## 3. Detection of bands and recovery of compounds.

absorbing, the bands were detected directly by viewing under uv light. If not, detection was done in conjunction with the sulphuric acid "charring" method, used on an analytical chromatogram run on the same sample under similar conditions. First, the corresponding bands as seen under uv light were scored on both analytical and preparative plates, then the analytical plate was sprayed with the 30% sulphuric acid solution and heated at 110 C for a few minutes. The location of the desired bands on the preparative plate could thus be inferred from the R value of the spots on the analytical plate.

Recovery of the separated compounds was achieved by removing the silica gel in each band from the plate, collecting it in a small glass filter and eluting the compound from the silica gel with a polar solvent such as acetone or methanol. Evaporation of the solvent then left the purified compound.

#### Measurement and recording of spectra

Mass spectra were recorded on an AEI model MS-2, an AEI model MS-9 or an AEI model MS-50 mass spectrometer. The MS-9 was used to determine molecular formulas by high resolution mass spectrometry (hrms). The MS-50 provided the atomic composition of all fragment peaks in a spectrum. All mass spectra in this thesis are recorded as the intensity of each peak relative to the base peak taken as 100%.

Proton nuclear magnetic resonance (nmr) spectra were recorded on either a Perkin-Elmer R 32 90 MHz spectrometer or a Varian Associates HA-100 100 MHz spectrometer. Spectra of very small samples could be recorded on this latter instrument in the Pulse Fourier Transform mode usin the Digilab FTS/NMR 3 Data system. C nmr (cmr) spectra were recorded on a Bruker WP-60 instrument. Chemical shifts are given throughout this thesis as  $\delta$  in ppm, using tetramethylsilane as reference standard. Spin-spin coupling data were obtained by way of double irradiation experiments. Nmr coupling patterns are described as follows: s - singlet, d - doublet, t - triplet, q - quartet, m - unresolved multiplet, u - unresolved absorption band.

Infrared (ir) spectra were recorded on a Perkin-Elmer model 421 dual grating spectrophotometer.

Ultraviolet (uv) spectra were recorded on a Unicam SP 1700 spectrophotometer.

## III. RESULTS AND DISCUSSION

## Characterization and partial purification of crude cyafrin

During the course of this work portions of crude cyafrin from six consecutive growths were obtained and examined. The amount of crude cyafrin varied between 75 and 150 mg per liter of medium. Crude cyafrin has the appearance of a dark brown, very viscous, oily substance.

Initial experiments by Mr. R. Parker of these laboratories had shown that crude cyafrin could be partially purified by a four funnel counter-current distribution between water and diethyl ether. It is assumed that the result of this procedure is to extract any highly polar substances such as carbohydrates and glycerol into the aqueous phase. Glycerol is one of the constituents of the Brodie medium used for growing C. africands. The combined ether fractions of the counter-current distribution yielded after evaporation of the solvent a light brown foam. This foam usually constitute about half the weight of the initial oil and was used as the starting material for all following separations.

0

Later the counter-current distribution was replaced by a simpler procedure which involved partitioning the crude cyafrin between ether and water. Crude cyafrin

was dissolved in the minimum amount of equal volumes of ether and water and the layers separated. The ether was extracted with an equal volume of water, then the two aqueous layers combined and extracted twice with an equal volume of ether. All ether layers were combined and the ether evaporated. Any residual water was removed by freezedrying. The results of this procedure were as satisfactory as those of the counter-current distribution.

Liquid-solid chromatography over silica gel has proven to be the most effective method of the several separation methods examined for the separation of crude cy-9,11,12 athin . For that reason liquid-solid chromatography was also the first choice for the separation of cyafrin.

Thin-layer chromatography (tlc) was considered more attractive than column chromatography because, once some skill has been acquired in handling and preparing the plates and applying the samples, it is a fast and convenient method and less prone to mechanical problems. Therefore preparative tlc has been the method used throughout this work, both for the initial large scale separations and for later purifications.

Preliminary tests indicated that solvent system A gave the most satisfactory separations of crude cyafrin.

Routinely the multiple elution technique was applied; three

elutions were usually sufficient.

It was found that the composition of crude cyafrin varied considerably between different growths. This is not surprising as a similar variability has been found with C. helenae. It is thought that mutations occurring in the mycelium during its growth are responsible for this 10,12 variation in composition. Thus the actual amount of any one metabolite isolated from crude cyafrin could vary from virtually none to as much as 10% of the weight of the crude mixture. Sometimes a certain band on preparative tlc would contain mainly one compound; at other times the same band would contain a complex mixture of compounds. The following results can therefore be considered as representing only a general trend.

In general most of the material was found on the high-polarity (low R ) side of the chromatograms. Usually a large proportion (40 to 50%) had an R of 0.0 (solvent f system A, triple elution). This material was not further examined.

Three components showed up on the that an expectation peared to contain substantial amounts of material (as judged from their size and colour-intensity after charring with 30% sulphuric acid). Of these, component A, R 0.5 and component B, R 0.2 (solvent system A, triple elution) were

coloured intensely purple by the charring while the third, component C, R 0.4, charred yellowish brown. These three components were selected for further examination.

Most compounds isolated during the course of this investigation were obtained and handled as more or less colourless gums.

# Isolation and identification of cyannin A<sub>3</sub> - allocyathin B<sub>3</sub> mixture

Taube had found the major metabolites of C. helenae to be cyathin A and allocyathin B (9 and 10 respectively). He found compounds 9 and 10 to have identical R (silica gel, solvent system Benzene: Acetone: Acetic acid 70: 30: 1) but he managed to separate 9 and 10 by tlc on silver nitrate impregnated silica gel:

As cyathin A and allocyathin B were the 3 3 3 most abundant of all cyathins isolated from C. helenae, it was decided to use them as reference substances for comparison with any compounds found in crude cyafrin.

Cyathin A (sample provided by Dr. P. Singer 3 of these laboratories) has an R value on tlc (R 0.5, solvent system A, triple elution) identical with that of component A found in C. africanus. Use of other solvent systems such as C (double elution) did not show any differences between component A and cyathin A.

The mass spectrum of component A showed four apparent parent ions at m/e 334, 332, 318 and 316. The ions at m/e 318 and 316 were the most intense with an intensity ratio of approximately 3:4. Other prominent ions were found at m/e 191, 189, 175, 45 and 43 (base peak). Determination of the molecular formulas of the parent peaks by high resolution mass spectrometry gave the following results: m/e 316, C H O; m/e 318, C H O, which are the same as the molecular formulas of allocyathin B and cyathin A . Thus component A is a mixture of four compounds, two of which are present in major amounts. The molecular formulas of the two minor parent peaks were found to be m/e 332, C H O 20 28 4 m/e 334, C H O. 20 30 4

The nmr spectrum of component A was very un-

informative, its main features being very broad, completely unresolved bands around  $\delta$  1.0, 1.5 and 2.1. In addition it showed unresolved narrower bands near  $\delta$  4.5 and b.0. This behaviour is not unexpected for cyathin-like compounds and 9 has been explained in terms of a fast tautomeric equilibrium between a cyathin and its internal hemi-ketal tautomeric form (Part structures 11 and 12).

<u>11</u>

It was decided to attempt to separate compo7, p.44
nent A by a method originally suggested by Taube . The
procedure involves acetylation of the hydroxyl groups of the
compounds in the mixture followed by preparative tlc of the
acetylated mixture. If desired, the isolated pure acetylated
cyathins can be deacetylated under mild basic conditions in
methanol to give the pure cyathins. An additional advantage

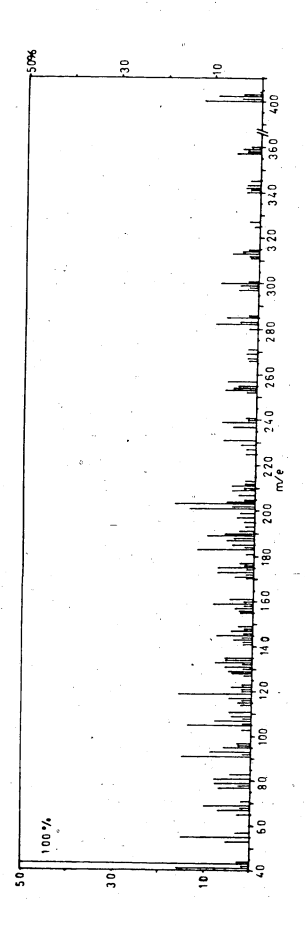
of this method is that the tautomeric equilibrium described above is not possible for acetylated cyathins, and it would therefore be expected that these derivatives would give more informative spectral data.

Acetylation of component A was achieved using pyridine and an excess of acetic anhydride overnight at room temperature. After concentration, the acetylated mixture was separated by prep. tlc (solvent system A) and the largest fraction, fraction A (at R 0.7) isolated and analyzed.

The mass spectrum of fraction A (see fig. 1) showed parent peaks at m/e 402 (hrms: C H O) and 400 24 34 5 (hrms: C H O) at an intensity ratio of about 3:4. Comparison of the spectrum of fraction A with the mass spectra of 0,0-diacetylcyathin A and 0,0-diacetylallocyathin B measured by Taube shows that fraction A is a 3:4 mixture of these two compounds.

In addition, the nmr spectrum of fraction A in CDC1 (see fig. 2) can be fully explained if fraction A is assumed to be composed of 0,0-diacetylcyathin A and 3 0,0-diacetylallocyathin B. Compare tables I and II with 7,9 fig. 2 (tables also from data published by Taube ).

It was concluded on this basis that the two



and 0,0-diacetylallocyathin B . Mass spectrum of fraction A (0,0-diacetylcyathin A

Figure 2. Nmr spectrum of fraction A (0,0-diacetylcyathin A and 0,0-diacetylallocyathin B ) (CDCl ).

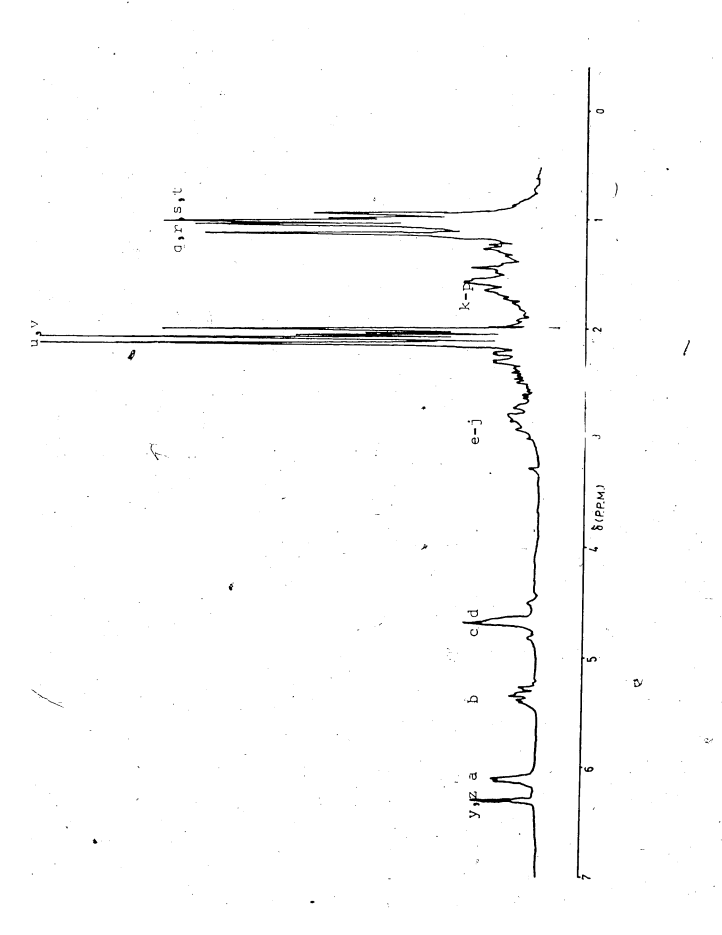


Table I. Nmr data of 0.0-diacetylcyathin  $A_3$  in  ${\rm CDCI}_3$ 

Assignment	13 11 15 18 5 10 2 10 1,7,8 (16,17) (19,20)
Coupled with signal	c,d,b d,a,b a,a,b i,t,b i,b,f e e e e
Coupling constant (Hz)	1.5,1.5,0.5 7,6,0.5,0.5 14,1.5,0.5 14,1.5,0.5 10,2.5 15,6,2.5 ?
Multiplicity	ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი
Number of protons	
Chemical shift (6)	6.13 5.36 4.74 4.63 2.85 2.78 2.53 approx. 2.1 approx. 1.9-1.1 1.15 1.06 1.00 0.99 2.12 2.04
Signal	«стюча х∪тя не до Фр - - - - - - - - -

For footnotes see page 26.

6

For footnotes see page 26.

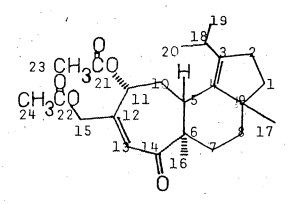
Footnotes to tables I and II.

Identification of signals by letters is as in ref. 7: letters are assigned alphabetically to signals belonging to protons on the basic cyathin structure in order of decreasing chemical shift; remaining letters are assigned to signals belonging to protons added on to the basic structure by derivation. This coding is also used in all following tables of nmr data.

For notation see general experimental.

The constants are listed in the same order as the signals in the following column.

Numbers refer to numbered positions in structure 13



major compounds present in component A are identical with cyathin A and allocyathin B. This work establishes that 3 3 C. africanus does indeed produce metabolites similar to those found in C. helenae.

## Isolation and proposed structure of cyafrin $A_{4}$

The next subject of investigation was component B (R 0.2, solvent system A, triple elution). After f initial isolation by prep. tlc using this system component B was found to be still contaminated with more polar material. Further purification was achieved by prep. tlc with solvent system D to give pure compound B as judged by tlc. On one occasion a sample was obtained crystalline, but usually compound B was handled as a gum.

The mass spectrum of compound B (see fig. 3) showed a parent peak at m/e 334 and other prominent peaks at m/e 203, 190 (base), 175 and 43. The material appeared to be essentially pure, but a small peak at m/e 332 suggested a minor impurity. Hrms gave a molecular formula of C H O for m/e 334 and showed the atomic composition of 20 30 4 the major fragment peak (m/e 190) to be C H . By analogy with the general scheme for naming cyathins compound B was called cyafrin A .

The nmr spectrum of cyafrin A (see fig. 4)

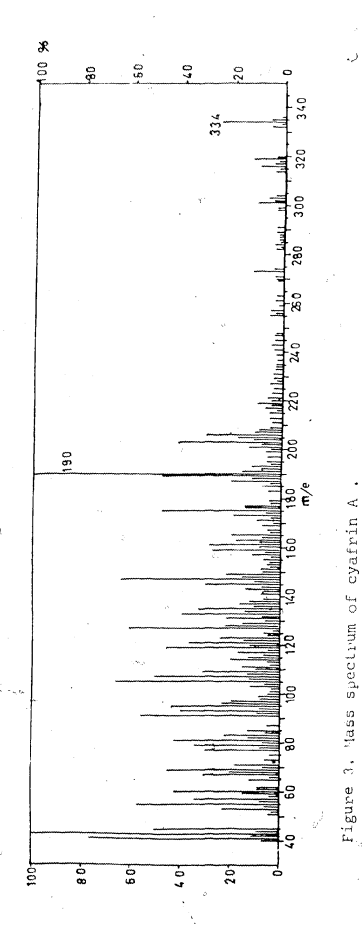
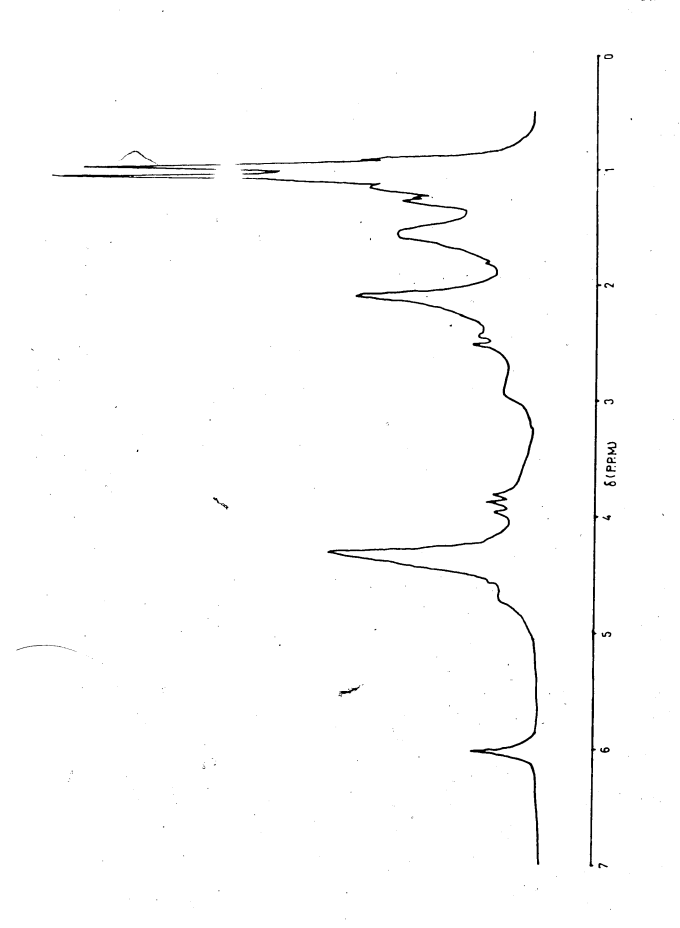


Figure 4. Nmr spectrum of cyafrin A (CDCl ).  $\frac{4}{3}$ 



turned out to be poorly resolved, as anticipated for cyathin-7, p.39 like compounds . For cyathins the poor resolution in the nmr is attributed to the ketone - hemiketal tautomeric equilibrium described earlier. The only distinguishable feature of the nmr spectrum of cyafrin A that is of interest is the triplet (J = 7.5 Hz) at  $\delta$  3.9, which is not present in the nmr spectra of cyathin A and allocyathin B.

The infrared spectrum of cyafrin A is shown in fig. 5 (in chloroform solution) and fig. 6 (neat). The second spectrum was obtained by applying a chloroform solution of cyafrin A to the face of the ir cell and allowing the chloroform to evaporate. Both spectra show the general features found in the spectra of cyathins. See for example The absorption at 3600cm the spectra of cyathin A in the solution spectrum may be attributed to free hydroxyl groups. In pure state all hydroxyl groups are hydrogen-bonded, as evidenced by the relatively more intense absorption in the neat spectrum (fig. 6), and the ketone. function is completely masked. This phenomenon is also observed in the if spectra of cyathins and is attributed to the formation of an internal hemiketal. It has been shown that cyathin A crystallizes exclusively in this form The remaining absorption of cyafrin A at 1650 cm possibly due to a number of C=C double bonds in the molecule. is due to residual chloroform. The sharp peak at 750 cm

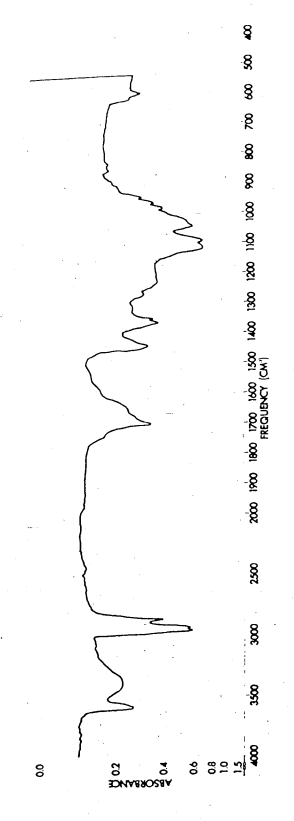


Figure 5. Ir spectrum of cyafrin A (CHCI

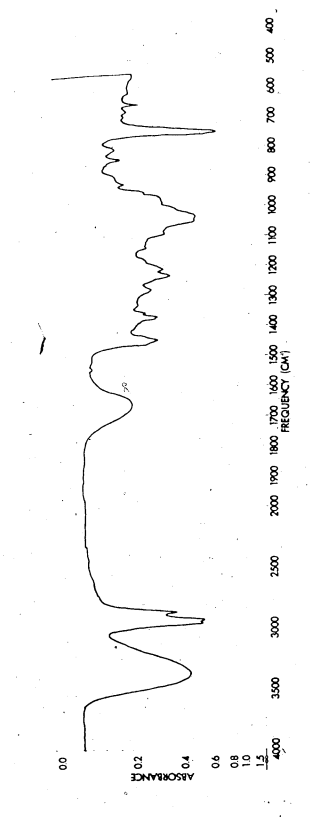


Figure 6. Ir spectrum of of frin A (neat).

It thus appears that cyafrin A possesses a cyathin-like structure containing a number of hydroxyl groups. To obtain information on the number of hydroxyl groups in cyafrin A it was acetylated. Acetylation of cyafrin A was achieved in the same manner as described for the cyathin A - allocyathin B mixture, using pyridine as 3 solvent and a large excess of acetic anhydride, overnight at room temperature. The crude product was purified by prep. tlc (solvent system A).

The mass spectrum of acetylated cyafrin A

(see fig. 7) shows an apparent parent peak at m/e 400,

hrms: C H O . This at first perhaps surprising result
24 32 5

may be explained if one assumes the formation of a triacetyl derivative. A triacetyl derivative (which would have a

molecular formula of C H O, m/e 460) can eliminate one
26 36 7

molecule of acetic acid in the mass spectrometer (high temperature, low pressure) if a proton is suitably positioned

on a carbon atom vicinal to that bearing the acetylated

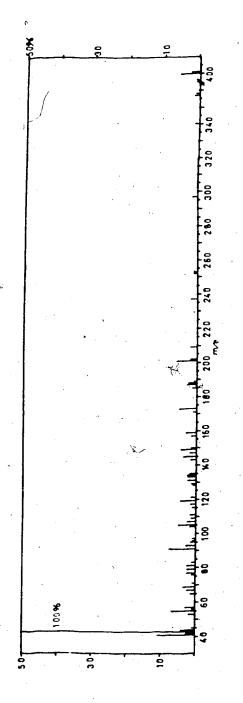
hydroxyl group involved. This process would result in the
fragment at m/e 400. The small peak at m/e 340 could be the
result of the loss of a second molecule of acetic acid. It
was thus tentatively assumed that cyafrin A had formed an
0,0,0-triacetyl derivative.

This assumption is confirmed by the nmr spec-

trum (see fig. 8). The three sharp singlets at 6 2.02, 2.07 and 2.12 are assigned to the protons of three acetyl groups. The nmr spectrum of 0,0,0-triacetylcyafrin A proved to be very instructive. Many of its features could be analyzed by virtue of its striking similarity to that of 0,0-diacetylcyathin A. Compare tables I and III. The data for 0,0,0-3 triacetylcyafrin A (table III) were obtained using a 100 MHz spectrometer; the information on couplings was obtained by means of spin-spin decoupling (double irradiation) experiments. Table I (0,0-diacetylcyathin A) was compiled from data obtained using a 220 MHz spectrometer, which accounts for its greater detail.

A comparison of table I and table III shows only three significant differences. Firstly, 0,0,0-triace-tylcyafrin A (table III) shows an extra signal at δ 4.89, which is coupled with some of the protons in the region of signals e to j. Secondly, the region between δ 1.9 and 1.1, which contains six protons in 0,0-diacetylcyathin A, contains only four in 0,0,0-triacetylcyafrin A. Thirdly, the region around δ 2.1 (signals u,v,w) shows three three-proton singlets in 0,0,0-triacetylcyafrin A instead of two (signals u,v) as observed in 0,0-diacetylcyathin A.

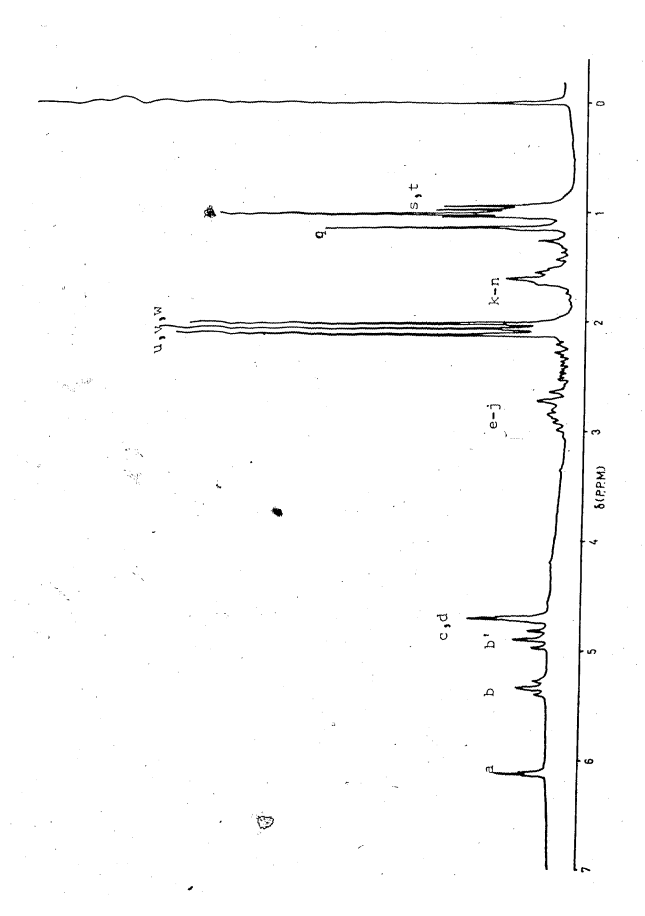
The conclusions one can draw from this comparison are: cyafrin A and cyathin A are likely to have



tgure /. Mass spectrum of 0,0,0-triacetylcyafrin A

•

Figure 8. Nmr spectrum of 0,0,0-triacetylcyafrin A (CDC1).



a 6.12 b 5.33 b' 4.89 c,d 4.70 e-j region e-j region c,d 4.70 e-j region k-n 1.9-1.1 t 5 a e-j region d 1.5 a b,b',s,t,?	Chemical shift (6) protons  Mull	
200	6.12 5.33 4.89 4.70 3.0-2.1 6 1.9-1.1 1.02 3.3 3.4 6.5 6-j reg 8 e-j reg 1.9-j reg 1.9-j reg 3.0-2.1 6 u 1.9-1.1 3.0-2.3 6 i 1.9-1.1 1.00 3.0-2.3 8 i 1.00	6.12 6.12 5.33 4.89 4.70 3.0-2.1 6.19-1.1 1.5 5.4 4.3 5.3 4.3 5.3 6.5 6.5 6.5 6.5 6.5 6.5 6.5 6.5 7.6 8 6.7 8 8 6.7 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 9 1.5 9 1.5 1.5 8 1.5 8 1.5 8 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0

. .

the same basic structure; of two protons absorbing in the methylene proton region for 0,0-diacetylcyathin A, one 3 has been replaced by a hydroxyl group (an acetoxyl group in 0,0,0-triacetylcyafrin A), the other by a proton which 4 absorbs in the same region as other protons geminal to an acetoxyl group.

If the first conclusion is correct some tentative structure assignments for cyafrin A may be arrived at. Looking at the structure of 0,0-diacetyleyathin A and at the assignment of proton signals in the molecule (see structure 13-) it is found that the molecule contains six methylene carbons that could in principle bear the extra acetoxyl group observed in 0,0,0-triacetylcyafrin A . They are carbons 1, 2, 7, 8, 10 and 15. Of these, C-10 can be ruled out: both protons on C-10 (signal g,j) are still present in the nmr spectrum of 0,0,0-triacetylcyafrin A, as is evidenced by the multiplicity of signal b (fig. 8, table III). C-15 is not a possible site as the nmr signal c,d (fig. 8) of 0,0,0-triacetylcyafrin A contains two protons. It seems then that cyafrin A could have the same skeleton as cyathin A, but would bear an extra oxygen on one of carbons 1, 2, 7 and 8.

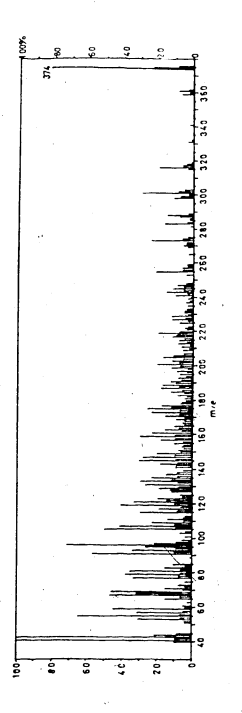
To find further support for the tentative structure assignment and to obtain further information about the position of the extra oxygen function we decided to prepare

a second derivative of cyafrin A. An attractive possibility presented itself at once. It has been found that cyathin A 12, p.102 3 readily forms an 0,0-isopropylidene derivative , which results in the formation of an extra six-membered ring. If cyafrin A does possess a similar basic skeleton it should form this derivative without difficulty, as the extra hydroxyl group is not likely (see above) to be in a position to interfere with the formation of an isopropylidene derivative. Another reason for preparing this derivative was this: a chanical conversion of cyafrin A to one of the known cyathins would prove our tentative structure assignment for cyafrin A. Selective protection of two of the hydroxyl groups of cyafrin A as the 0,0-isopropylidene derivative would leave the third hydroxyl free for chemical transformations.

The 0,0-isopropylidene derivative (cyafrin A up acetonide) was prepared by dissolving cyafrin A in 2,2-dimethoxypropane. The reaction was catalyzed with p-toluenesulphonic acid. Direct application of the reaction mixture to a prep. tlc plate and elution (solvent system A) resulted in the isolation of pure (by tlc) cyafrin A acetonide.

The mass spectrum of cyafrin A acetonide (see fig. 9) shows a strong parent peak at m/e 374 (hrms:

C H O) as expected for a mono-0,0-isopropylidene deriva23 34 4



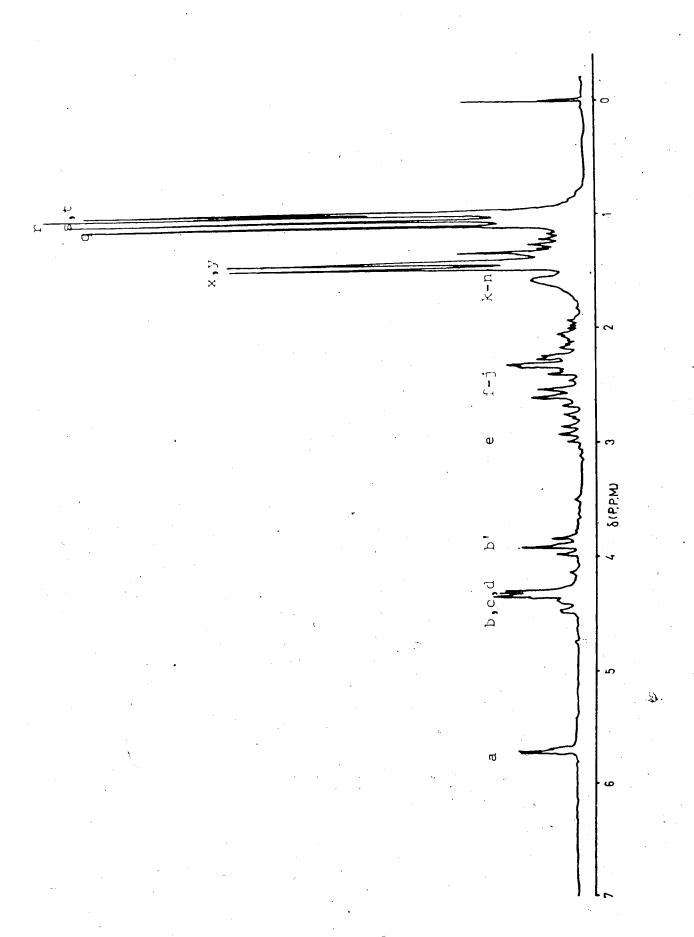
tive of a C H O compound, and no evidence of impurities.

The nmr spectrum of cyafrin A acetonide (see fig. 10) confirms that it is an 0,0-isopropylidene derivative: the two three proton single's around  $\delta$  1.45 are attributed to the newly introduced methyl groups.

As was the case with the spectrum of the triacetyl derivative of cyafrin A the nmr spectrum of cyafrin A acetonide can be analyzed by comparison with that of Cyathin A acetonide. This compound was first prepared by Mercer, and table V is from data collected by him Comparison of the data in this table with the nmr data for acetonide (table IV) shows many similarities. The data in the last two columns of table IV were obtained by means of spin-spin decoupling (double irradiation) ex-Periments. The differences between the data in these two tables are very instructive. The spectrum of cyafrin A acetogide shows the extra signal b' at 8 3.9, and there appear to be two protons less in the  $\delta$  1.2-1.8 region. This confirms Our previous conclusions: signal b' is attributable to a proton on a carbon that also bears a hydroxyl group. Signal b' is coupled with two protons ( $\delta$  2.3-2.5) by equal or almost $^\omega$ equal spin-spin coupling constants.

The question remained: which of the carbons 1,  $^2$ , 7 and 8 is the one bearing the hydroxyl group? The chemi-

Figure 10. Nmr spectrum of 0,0-isopropylidenecyafrin A (CDC1)



in CDC1 Table IV. Nmr data of 0,0-isopropylidenecyafrin A

Coupled with	b,c,d region a,? 2.3-2.5 region s,t	
Coupling constant (Hz)	small ? 7.5 % . 5	<b>υ</b>  
Multiplicity	ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი ი	
Number of protons	പെ പെ ⇔കുട	ოო
Chemical shift (6)	5.71 4.3-4.5 3.90 2.92 2.8-1.8 1.8-1.2 1.00 1.00	1.42
Signal	b,c,d b,c,d K-1 x,t	<b>&gt;</b>

in CDC1 Table V. Nmr data of 0,0-isopropylidenecyathin A

(-)

Signal	Chemical shift (8)	Number of protons	Multiplicity	Coupling constant (Hz)	a Assignment
ਰ	2,68	H		smal1	13
p'o'q	4.25-4.5	m -	n	c·	11,15
υ	2.92	7	<b>6 6</b>	6.5.	18
f-j	2.8-1.8	വ	בי	۷۰.	2,5,10
k-p	1.8-1.1	9	· p	· ·	1,7,8
<b>5</b>	1.10	က	ဟ		(16,
ង	1.06	က	<b>ග</b>		17)
ຜ	1.02	m	. °O	0.5	(19,
4	66.0	က	ט	ယ မ	20)
×	1.47	က	· ω		(22.
>	1.40	ന	Ø		23)
				20	

707	23/20 13/2	1,20	15 13 14	<b>3</b> 0	14
	22	``	· .	,	

Numbers refer to numbered positions in structure 14 (0,0-isopropylidene-

cyathin A .

cal shifts of the methylene protons which are coupled with the proton responsible for signal b', as we have found them in the spectra of the triacetyl- and the acetonide derivatives (2.1-3.0 and 2.3-2.5 ppm respectively, see tables III and IV) make positions 7 and 8 unlikely to be the ones: the presence of a hydroxyl group on an adjacent carbon would not be expected to deshield these protons appreciably from, e.g., the shift of 1.44 ppm found for the methylene protons in cyclohexane The literature was searched for nmr data on suitable model compounds to make a decision between positions 1 and 2 possible. No data on 2-cyclopenten-1-ol or similar compounds were found, but data on some 3-cyclopenten-1-ol systems are available. The relevant chemical shifts are indicated on the corresponding structures 15 (from ref. 19) and 16 (from ref. 20). These values correspond closely to our values: signal b' is found at δ 4.89 in triacetylcyafrin A and at & 3.90 in cyafrin A acetonide. The methylene protons resonate at  $\delta$  2.1-3.0 and  $\delta$  2.3-2.5

16

in these two derivatives respectively. It is not likely that the protons in 2-cyclopenten-1-ol systems would have similar chemical shifts, as is shown by the large difference between the two sets of methylene protons in cyclopentene itself (structure 17, from ref. 21).

<u>17</u>

It was therefore tentatively assumed that the third hydroxyl group in cyafrin A is positioned on C-1.

oxidation of the free hydroxyl group in 0,0isopropylidenecyafrin A offered a second method of ascertaining the position of the third hydroxyl group of cyafrin
A. Oxidation of 0,0-isopropylidenecyafrin A was achieved
using "Jones" conditions (see detailed experimental) using
chromic acid. After filtration and separation of the product mixture by prep. tlc (solvent system A) a small quantity of chromatographically pure material was isolated. Mass
spectral analysis of this material indicated that it was the
expected oxidation product: a parent peak was found at m/e
372 (which corresponds to the molecular weight of the starting material minus two protons).

The infrared spectrum of the oxidation pro-1
duct showed carbonyl absorption at 1735 cm. This is consistent with the presence of a non-conjugated carbonyl
group in a five-membered ring.

The nmr sp frum of the oxidation product showed as new feature a two proton singlet at  $\delta$  3.00, while the triplet found in the spectrum of cyafrin A acetonide at  $\delta$  3.90 had disappeared. The two proton singlet may be assigned to the two protons adjacent to the newly introduced carbonyl group. The fact that these are magnetically equivalent is an indication that they may be positioned on a five-membered ring. In addition, the chemical shift of these

two protons is such ( $\delta$  3.00) that it can only be accommodated in the cyathin skeleton by assuming them to be positioned on C-2, which is flanked both by the new carbonyl group and by the C=C double bond. This confirms that the extra hydroxyl group in cyafrin A is positioned on C-1.

. We can now arrive at a more precise formulation (structure  $\underline{18}$ ) of cyafrin A .

18

The only problem that remains to be solved is that of the stereochemistry of the C-l hydroxyl group, i.e., is the hydroxyl group orientated cis to the methyl group on C-9 or trans to it?

It is known that when nmr spectra are recorded in pyridine the chemical shifts of protons which are

positioned close to carbonyl or hydroxyl groups may change considerably when compared to their shifts in non-aromatic solvents such as chloroform. It is assumed that the pyridine complexes with these functional groups, thus changing the magnetic environment of nearby protons. In the case where a methyl group is situated on a carbon vicinal to a carbon bearing a hydroxyl group, a definite relationship has been between the dihedral angle (formed by the found to exist C-OH bond and the C-CH bond) and the difference in chemical shift of the methyl protons when measured first in chloroform and then in pyridine solution. Thus for a synclinal relationship between hydroxyl and methyl groups, a downfield shift of 0.20 PPM or larger may be expected. For an antiperiplanar relationship the shift will be 0.10 - 0.12 PPM or less. By recording the nmr-spectrum of 0,0-isopropylidenecyafrin A in pyridine-d and noting how the chemical shifts of the methyl groups change compared to the shifts in CDCl (as given in table IV) it was hoped that the orientation of the hydroxyl group at C-1 could be deduced. Table VI shows how the shifts of the methyl groups compare for the chloroform and pyridine spectra. The doublet signals are due to the protons on the isopropyl group. Two interpretations are possible for the tertiary methyl signals. Either both signals have shifted by 0.21 PPM (from  $\delta$  1.00 to  $\delta$  1.21 and from  $\delta$  1.09 to  $\delta$  1.30 respectively) or one has shifted by 0.30 PPM (from  $\delta$  1.00 to  $\delta$  1.30) and the

Table VI. Chemical shifts of methyl protons in 0,0-isopropylidenecyafrin A

Shift in CDCl (δ) Multiplicity 3	Shift in C D N (8) M	ultiplicity
1.09	1.30	S
1.02	1.21	S
1.00	1.01	d
	0.91	d

Integrates to six protons.

other by 0.12 PPM (from δ 1.09 to δ 1.21). The first of these two interpretations is the more attractive since it eliminates the need to establish which signal corresponds to the C-17 methyl group and which to the C-16 methyl group. It is not unusual that the C-16 methyl group shifts as well as the C-17 methyl group, since it is near the carbonyl group. This would then lead to the conclusion that the hydroxyl group at C-1 must be orientated cis to the vicinal methyl group.

The second possible interpretation must not be neglected. Assuming that one methyl signal had shifted by 0.12 PPM and the other by 0.30 PPM it becomes necessary to establish which signal belongs to which methyl group. To determine this the spectrum of 0,0-isopropylidenecyathin A

(prepared by the method of Mercer ) was recorded in pyridine and compared with that recorded in chloroform (see table V). Cyathin A is not functionalized in the five-membered ring, so the shift of the C-17 methyl protons is not expected to be influenced appreciably by the change of solvent. If we see a shift of one of the methyl signals then that shift must be associated with the C-16 methyl group. The nmr spectral results are given in table VII. The isopro-

Table VII. Chemical shifts of methyl protons in 0,0-isopropylidenecyathin A

Shift in CDCl (8)	Multiplicity	Shift in C D N (δ) 5 5	Multiplicity
1.10	s	1.18	S
1.06	S	1.06	s
1.02	d	0.96	đ
0.99	đ	0.92	ď

pyl signals are readily recognized by their multiplicity. The chemical shift of the C-17 methyl protons should not change markedly and must be that of the signal at  $\delta$  1.06. The signal of the C-16 methyl protons has shifted from  $\delta$  1.10 to  $\delta$  1.18, i.e., 0.08 PPM downfield. Thus in cyafrin A acetonide the signal which shifts by 0.12 PPM must be the signal due to the C-17 methyl protons, and that which

shifts 0.30 PPM must be from the C-16 methyl protons. Hence the hydroxyl group at C-1 must be orientated cis to the C-17 methyl group.

It is possible to confirm this stereochemical 23 assignment using C-nmr. Recently Djerassi et al measured 13 C-nmr spectra of a number of steroids in which a hydroxyl group and a methyl group were positioned on adjacent carbons in a five-membered ring. They noted that if the relation-ship between hydroxyl and methyl group was antiperiplanar, the chemical shift of the methyl carbon was only 0.4 PPM upfield from the case where the hydroxyl group was absent. If the relationship between them was synclinal, the methyl group was shifted upfield about 6 PPM.

The C-nmr spectra of several known cyathins, including 0,0-diacetylcyathin A , have recently been measured in these laboratories, and the signals assigned to specific 24 carbon atoms .

A large quantity (45 mg) of 0,0,0-triacetyl13
cyafrin A was prepared and a C-nmr spectrum was obtained
4
on this material. The shifts were compared with those of
0,0-diacetylcyathin A (see table VIII). Many signals have
3
similar shifts to those of 0,0-diacetylcyathin A . These
3
have tentatively been assigned to the corresponding carbons.

J

Table VIII. C-nmr data of 0,0-diacetylcyathin A and 3 0,0,0-triacetylcyafrin A in CDC1.

0,0-diacetylo	yathin A 3	0,0,0-triacet	ylcyafrin A
Chemical shift (PPM)	a Assignment	Chemical b	Chemical c shift (PPM)
208.7 170.1 169.8 143.4 140.7 135.5 127.4 72.0 64.7 54.3 49.3 38.7	14 (21, 22) 12 (3, 4) 13 11 15 6 9	143.5 138.0 134.4 127.3 71.8 64.7 53.6 50.4 39.6	80.8 34.4 21.9 17.1
38.0 36.2 34.7 31.7 28.2 27.2 24.0 21.7 21.7 20.8 20.8 14.8	(1, 7) 8 10 2 18 17 (19, (19, 20) (23, 24)	35.4 34.9 31.7 27.1 21.1 20.8 20.8 15.0	

Numbers refer to carbons numbered as in structure 13, p.26.

only the shifts of signals closely corresponding to signal in the first column are listed here.

The shifts of signals not listed in the previous column are listed here.

Several major differences are also apparent. First of all, only 22 signals were found for 0,0,0-triacetylcyafrin A , although the compound contains 26 carbons. At the time the spectrum was recorded samples larger than 45 mg were usually required for good results. The missing signals are those of the carbonyl carbons (C-14 and the acetyl carbons), which are relatively weak signals. The signal assigned to C-l in 0,0-diacetylcyathin A has apparently shifted to &80.8, close to the signal for C-ll, the other methine carbon bearing an acetoxyl group in 0,0,0-triacetylcyaf~in A .\* signal for C-2 in 0,0-diacetylcyathin A has shifted from δ 28.2 to δ 34.3 in 0,0,0-tr. yafrin A , due to the influence of the acetoxyl group at M-1. In the 0,0,0-triacetylcyafrin A a ne or gnal is found in the me-"hyl region at & 21.9: this must be the methyl carbon of the acetoxyl group. Finally the signal assigned to C-l/ in in 0,0-diacetylcyathin A has shifted upfield from δ 24.0 to  $\delta$  17.1, <u>i.e.</u>, by 6.9 PPM, in 0,0,0-triacetylcyafrin A which compares with a shift of about 6 PPM in the case of a synclinal relationship observed by Djerassi , and which is a typical  $\gamma$ -gauche effect.

This confirms our earlier finding that the hydroxyl group at C-l in cyafrin A must be orientated cis to C-l7. Structure 19 is therefore proposed for cyafrin A

<u>19</u>

## Correlation of cyafrin Au with allocyathin B3

Abundant s actral evidence has been presented to support structure  $\underline{19}$  as that of cyafrin A. The finding of cyathin A and allocyathin B as major metabolites 3 of C. africanus shows that the structure for cyafrin A<sub>4</sub> is biogenetically sonable. In order to prove conclusively that cyafrin A<sub>4</sub> has structure  $\underline{19}$ , a chemical correlation of cyafrin A<sub>4</sub> with one of the known cyathins was desirable.

We proposed to do this (as illustrated in scheme I) by dehydration of 0,0-isopropylidenecyafrin A to give 0,0-isopropylideneallocyathin B, a compound which has been prepared from allocyathin B and described by Mercer.

Many reagents are known which can effect dehydration. However, many of them employ acidic conditions

Scheme I. Proposed conversion of 0,0-isopropylidenecyafrin A to 0,0-isopropylideneallocyathin B.

which should be avoided in this case because the 0,0-isopropylidene protecting group is acid labile. A procedure that, employs basic conditions developed by Von Rudloff used successfully for the dehydration of various natural products and derivatives . This method consists of heating the alcohol in the presence of alumina impregnated with 1 o 2% of pyridine. We decided to replace pyridine by colliline to minimize loss of the reagent by evaporation. A small amount of cyafrin A acetonide was heated with this catalyst at 150 C for 30 minutes (these conditions are much milder than the ones suggested by Von Rudloff). The products were washed from the catalyst with methylene chloride and then methanol. The product mixture was examined by tlc, using authentic allocyathin B acetonide as reference substance. However, no trace of the desired product nor any starting material could be detected in the product. Discouraged by these findings we decided to look for less vigorous dehydration methods.

Another well known dehydrating agent is thionyl chloride, which may be used in combination with a base such as pyridine. When we first attempted this reagent, pyridine was used as solvent and a large excess of thionyl chloride was added. The reaction was allowed to proceed at 0 0 C for three hours. After this time part of the reaction mixture was worked up and examined by tlc. It appeared that some of the starting material had disappeared, but little else could be detected. After a further four hours reaction at room temperature the product was again examined. The result this time was that most of the starting material had disappeared. The bulk of the reaction mixture now appeared to have an R of 0.0 (solvent system A) but wery faint f spot was just discernable at the R expected of allocyathin B acetonide (R 0.8).

Dehydration was then attempted using methylene chloride as solvent, with excess pyridine and thionyl chloride for 24 hours at room temperature. Examination of an aliquot of the reaction mixture showed very little change after 6 hours. After 24 hours tlc indicated a similar result as before: very polar material and a faint spot around R 0.8.

It was decided to use acetone as solvent (as this should insure the retention of the acetonide protecting group) and smaller amounts of reagents. The reaction was monitored by direct application of aliquots of the reaction mixture to the plates. This method proved useful: using 10 equivalents of freshly distilled thionyl chloride and 30 equivalents of pyridine in dry acetone at 0°C and monitoring the reaction by the (solvent system A) it was found that after 50 minutes the reaction was complete. The the of the reaction product showed not only highly polar material, but also a substantial amount of material with R 0.8, which is the R value of allocyathin B acetonide. The material at R 0.8 was isolated by prep. the. The yield was somewhat f less than 2 mg from 3.5 mg of starting material.

The mass spectrum of the dehydration product (see fig. 11) appeared promising. The apparent parent peak at m/e 356 is that expected for allocyathin B acetonide. In fact, this spectrum was identical with the mass spectrum of allocyathin B acetonide recorded by Mercer.

The nmr spectrum of the "dehydration profluct" (see fig. 12) was, however, disappointing. This spectrum (recorded at 60 MHz on a small quantity of material) showed all the features expected of a cyathin acetonide: two methyl signals around  $\delta$  1.45, two more just above  $\delta$  1.00 and ab-

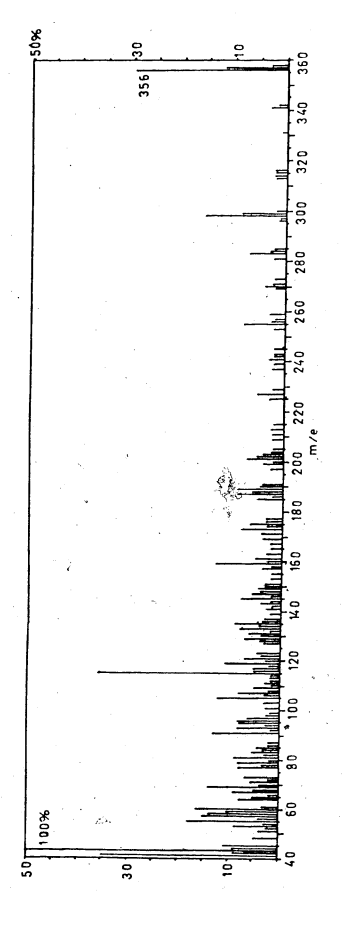
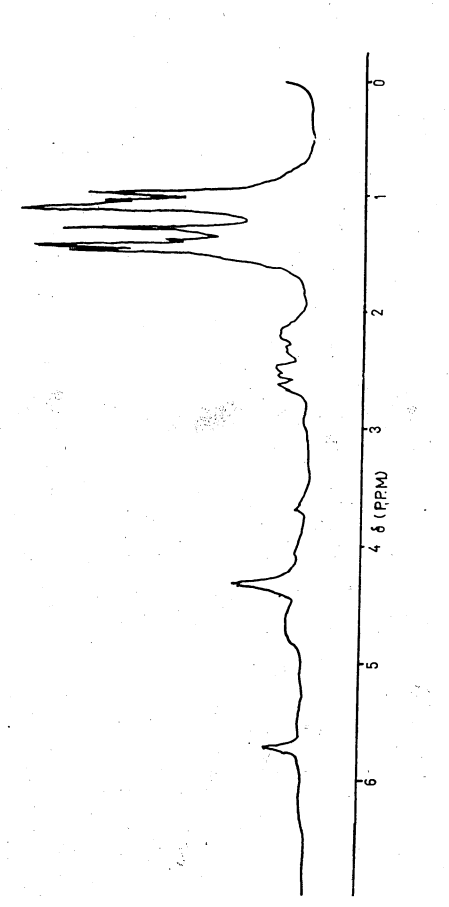


Figure 11. Mass spectrum of the dehydration product of cyafrin A



acetonide (CDC1 Figure 12. Nmr spectrum of the "dehydration product" of cyafrin A

sorption around  $\delta$ 1.0 that indicate the presence of an isopropyl group. Also, fairly sharp signals are found at  $\delta$ 4.3 and 5.7 (compare signals b,c,d and a in tables IV and V). However, allocyathin B acetonide has two olefinic protons on C-1 and C-2 that are known to resonate around  $\delta$ 6.3, and the reaction product (fig. 12) does not show signals in this region. In addition, fig. 13 shows a broad signal around  $\delta$ 4.8 which is not present in the spectrum of allocyathin B acetonide.

The uv spectrum of the material in hexane confirmed that the material is not identical with allocyathin. B acetonide; it showed absorption maxima at 231 nm ( $\epsilon \sim 5000$ ) and 330 nm ( $\epsilon \sim 100$ ). These are the normal values for the  $\alpha,\beta$ -unsaturated ketone found in all cyathins. No trace w found of a shoulder at 256 nm, as would be expected for the 7,9 conjugated diene present in allocyathin B acetonide .

It is known than when thionyl chloride is used as dehydrating agent, sometimes the reaction that takes place is substitution instead of elimination. That is, in our case, we could have obtained 1-chlorocyathin A acetonide (see structure 20) instead of allocyathin B acetonide. This would account for the evidence described: 1-chlorocyathin A acetonide would not show diene absorption in the ultraviolet region, nor would it show signals at & 6.3 in the nmr spectrum, but it could well show a signal around & 4.8, due to

20

the proton remaining on C-1. The mass spectrum may be explained by assuming that 1-chlorocyathin A acet ide, when the acet ide, when the spectrometer of the spectrometer of the corresponding radical ion.

B acetonide or the corresponding radical ion.

At this point we recalled that the mass spectrum of 0,0,0-triacetylcyafrin A (see fig. 7) showed the highest mass peak at m/e 400 instead of 460 as expected, which we attributed to the molecule losing a molecule of acetic acid in the mass spectrometer. If this molecule of acetic acid originated from the acetoxyl group on C-l and a proton on C-2, this would result in the formation of 0,0-diacetylallocyathin B. Comparison of fig. 7 with the mass spectrum of allocyathin B diacetate as recorded by Taube showed them to be identical.

The mass spectral correlation strongly suggests that cyafrin A does have the same basic skeleton as allocyathin B. However, the direct correlation has not been achieved, and the possibility that there are stereochemical differences (relative or absolute) has not been eliminated.

Further attempts to obtain more of what is assumed to be 1-chlorocyathin A acetonide were not undertaken because of a lack of starting material: during the later stages of this work the amount of cyafrin A isolated from growths of C. africanus had decreased. The area on prep. tlc that earlier contained pure cyafrin A had now begun to contain increasing amounts of other compounds, which proved very difficult to remove. Presumably mutations had occurred in the fungus and it will be necessary to obtain a new stock culture before the direct correlation can be completed.

## Isolation and propose structure of cyafrin Bu

We now turned our attention to component C found on tlc at R 0.4 (solvent system A, triple elution).

Mass spectral analysis of component C, which was isolated by prep. tlc (solvent system A, triple elution), in most ses showed several apparent parent ions. The major one by far was at m/e 332 (hrms: C H O<sub>D</sub>). Other high mass 20 28.4

peaks appeared at m/e 348, 334 and 314 (hrms: C H O, 20 28 5
C H O and C H O respectively), with further prominent 20 30 4 20 26 3
peaks at m/e 219 (C H O) and 205 (C H O). This material 15 23 14 21
thus appeared to be composed mainly of a compound with molecular formula C H O, which according to our system for 20 28 4
naming cyafrins was called cyafrin B. It was, however, contaminated with small amounts of several other compounds.

Cyafrin B was purified by prep. tlc, using solvent system C, quadruple elut ... After this purification the mass spectrum looked very much cleaner (fig. The fragments m/e 205, C H O and m/e 219, C H O obviously 14 21 15 23 do originate from cyafrin B.

The ir spectra of cyafrin B, recorded in 4

CHC1 solution and neat (fig. 14 and 15 respectively) proved 3 interesting (the latter spectrum was obtained by evaporating the solvent from a solution of cyafrin B which had been applied to the surface of an ir cell). The solution -I spectrum shows free hydroxyl (3600 cm ), hydrogen bonded -1 hydroxyl (3400 cm ) and strong carbonyl (1685 cm ) absorptions. As was the case with cyafrin A the free hydroxyl absorption disappears in the neat spectrum. The carbonyl absorption is also less intense. This phenomenon is often observed for cyathin-like compounds.

The uv spectrum of cyafrin B shows the ge-

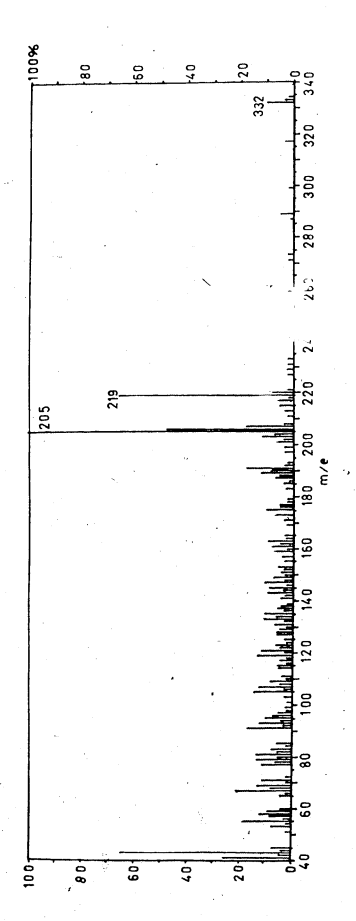


Figure 13. Mass spectrum of cyafrin B .

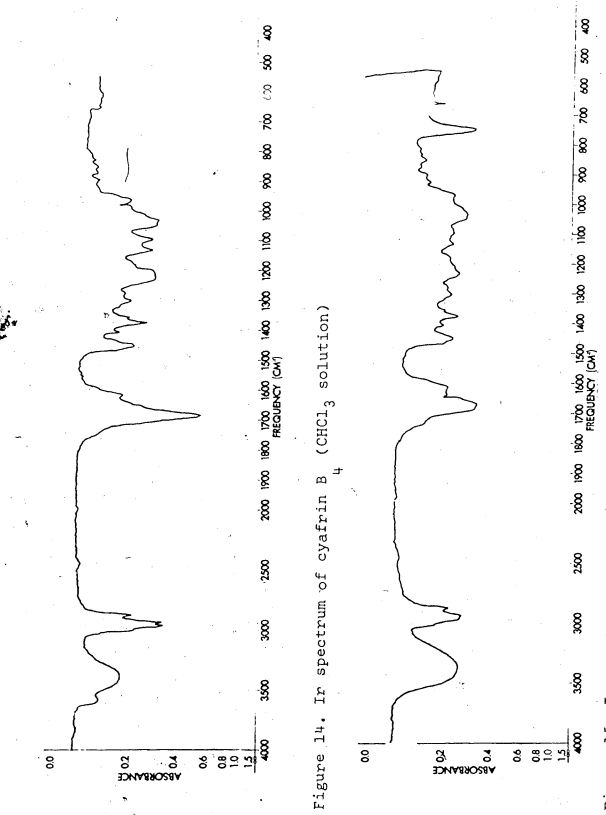


Figure 15. Ir spectrum of cyafrin  $B_{\mu}$  (neat)

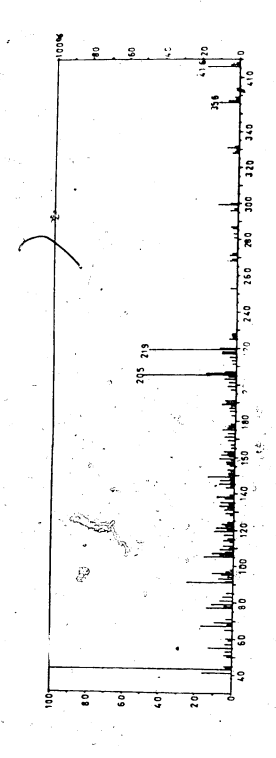
neral features of all cyathin-like compounds: a maxim 231 nm ( $\epsilon$  ~7000) and a shoulder around 305 nm ( $\epsilon$  ~100). Both may be attributed to the  $\alpha$ , $\beta$ -unsaturated ketone function present in the cyathins.

In view of the small amount of material present, an nmr spectrum of cyafrin B was not recorded since it would not be expected to yield much information. Instead an acetylated derivative of cyafrin B was prepared, since this would give information about the number of hydroxyl groups in the molecule and enable us to record well resolved spectra. Cyafrin B was acetylated in the manner described previously for cyafrin A. The product was isolated by prepared (solvent system A) and further purified by the same method (solvent system B, double elution).

The mass spectrum of the detylated product (see fig. 16) has a parent peak at m/e 416 (C H O). This 24 32 6 indicates that the product is a diacetyl derivative of cyafrin B. The peak at m/e 356 is attributed to loss of one molecule of acetic acid from the parent ion. The peaks at m/e 205 and 219 are still present, indicating that a major portion of the molecule has not been affected by the acetylation.

The infrared spectrum of 0,0-diacetylcyafrin

B (see fig. 17) confirms the presence of the acetoxyl



and i

Figure 16. Mass spectrum of 0,04diacetyloyafrin B

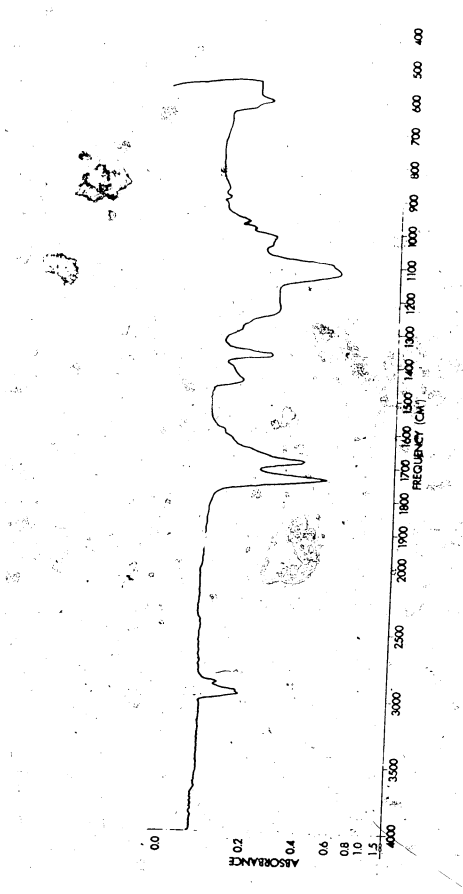
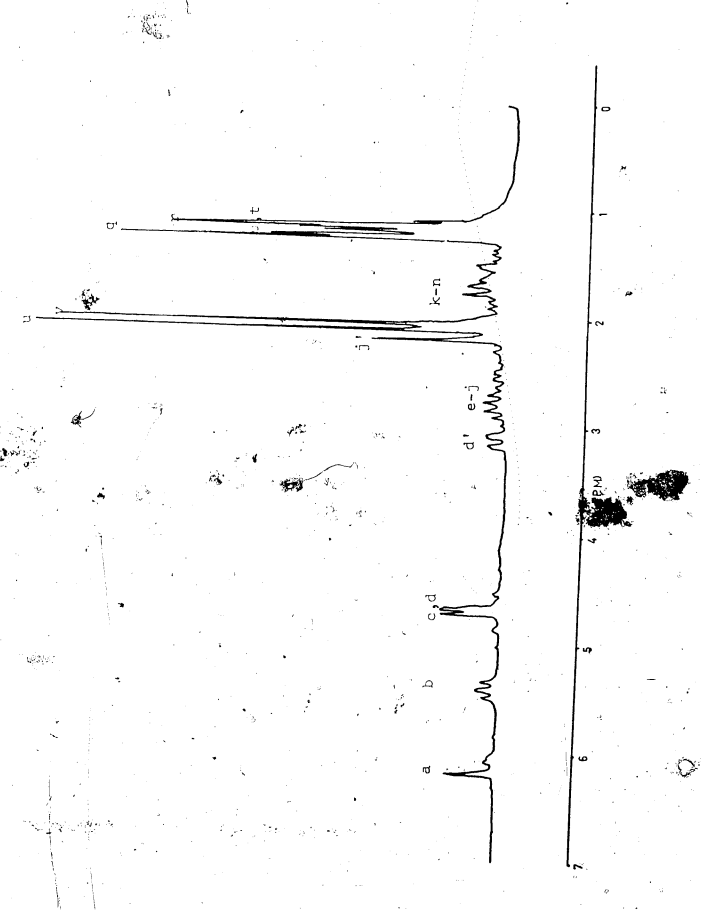


Figure 17. In spectrum of 0,0-diacetylcyafrin B

groups (carbonyl absorption at 1740 cm $^{-1}$ ). The absorption at 1685 cm $^{-1}$  is attributed to the  $\alpha$ , $\beta$ -unsaturated ketone function common to all cyathins. The absence of absorption around 3500 cm $^{-1}$  shows that all hydroxyl groups in the molecule have been acetylated.

The nmr spectrum of 0,0-diacetyloyafrin B (see fig. 18) has some interesting features. First, it confirms the presence of two retains by the appeareance of two three-proton singlets at 6 2.06, and 2.12. Second, the overall appearance of the spectrum is very similar to that of other acetyl derivatives of cyathins. Comparison of the data for this spectrum (table IX) with, for example, table I is very instructive. The signals a, b, c, d, u and v appear in the spectrum of 0,0-diacetylcyafrin B (table IX) virtually unchanged from those observed in the spectrum of 0,0diacetylcyathin A (table I) region between 8 2.25 and 3.0 of the nmr of 0,0-diacetylcyafrin B contains signals for no more than three protons, compared to six in 0,0-diacetylcyathin A (remembering that the molecular formula indicates two hydrogens less than for cyathin for cyafrin B A ). The "missing" signal in the nmr of 0,0-diacetylcyafrin has perhaps shifted to become signal d' at  $\delta$  3.13. The , sharp signal j' at  $\delta$  2.21 is difficult to account for at this stage: possibly it could originate from an impurity in the sample. Signals corresponding to the signals q, r, s and t

Figure 18. Mmr spectrum of 0,0-diacety\_cyafrin B



	_	Coupling constant (Hz)	L 0	1,1 1,1	10,1.5	¢.		,
afrin B in CDCl ﴿		Number of Multiplicity	1 1	р <b>,</b> р,	d,d	about a	<b>10</b> 10 10	, w c
Table IX. Nmr data of 0,0-diacetylcyafrin		Signal Chemical shift (6)	6.18 b 5.41	14.75 14.65 3.13	a-j 2 3 0-2.25	1.9-1.3 1.28	1.23	2.12 2.06

in the nmr of 0,0-d act ylc athin A are found to have shifted considerably to the while ield side of the spectrum in 0,0-diacetylcyafrin B. From these data a few conclusions can be drawn: of four oxygen atoms present in cyafrin B two exist as hydroxyl groups and a third forms part of an  $\alpha$ ,  $\beta$ -unsaturated ketone group; the seven-membered ring containing this  $\alpha$ ,  $\beta$ -unsaturated ketone and bearing the two hydroxyl groups in cyathin A is present unchanged in cyafrin B as evidenced by the signals a, b, c and d in the nmr spectrum of the diacetyl derivative; the molecule, like other cyathins, contains two tertiary methyl groups and an isopropyl group.

Other inferences may be made: cyafrin B may have the same basic skeleton as all cyathins; the fourth oxygen atom is present in a functional group which considerably influences the chemical shifts of many protons in the molecule (signals d', q, r, s and t in the nmr spectrum are all downfield compared to their "normal" positions).

The problem is now to establish the nature of the fourth oxygen function. This oxygen atom does not exist as a third hydroxyl group; the ir spectrum of the diacetyl derivative proves this. The  $\alpha$  , $\beta$  -unsaturated kerone group present in all cyathins very often cannot be detected because it can be "masked" by virtue of the internal ketonehemiketal equilibrium. We must therefore consider the possi-

bility of a second carbonyl group in cyafrin B, similarly undetected. It is also possible that the fourth oxygen is present as an ether function, e.g. as an epoxide. One of the metabolites of C. helenae, neoallocyathin A, has been shown to contain an epoxide function (see structure 7).

Taube first prepared methyl ketal derivatives of cyathin A and allocyathin B by treatment with methanol containing hydrogen chloride, as illustrated in scheme II.

Scheme Preparation of methyl ketal derivatives of cyathins

The preparation of such a derivative could assist us in ascertaining whether or not cyafrin B contains a second carbonyl group, since in this derivative the  $\alpha,\beta$ -unsaturated ketone is ketalized. If a further carbonyl group is present, this may now be observed in the infrared

spectrum.

The methyl ketal of cyafrin B was prepared 7,p.132;9 4
by the method of Taube . The product was isolated by prep. tlc of the reaction mixture (solvent system A, double elution, R 0.4).



Mass spectroscopy (see fig. 19) indicates

that the product is the desired cyafrin B methyl ketal

(parent peak: m/e 346, hrms: C H O). The mass spectrum

21 30 4

shows a very intense peak at m/e 141 (hrms: C H O) which

7 9 3

is common to methyl ketal derivatives of all cyathins. Its

formation has been rationalized as illustrated in scheme

7, p.72; 9

III

Scheme III. Fragmentation of a cyathin methyl ketal

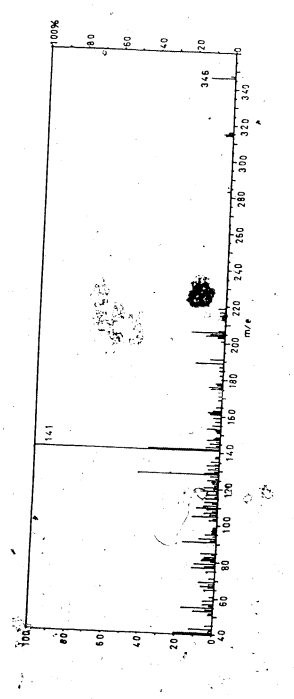
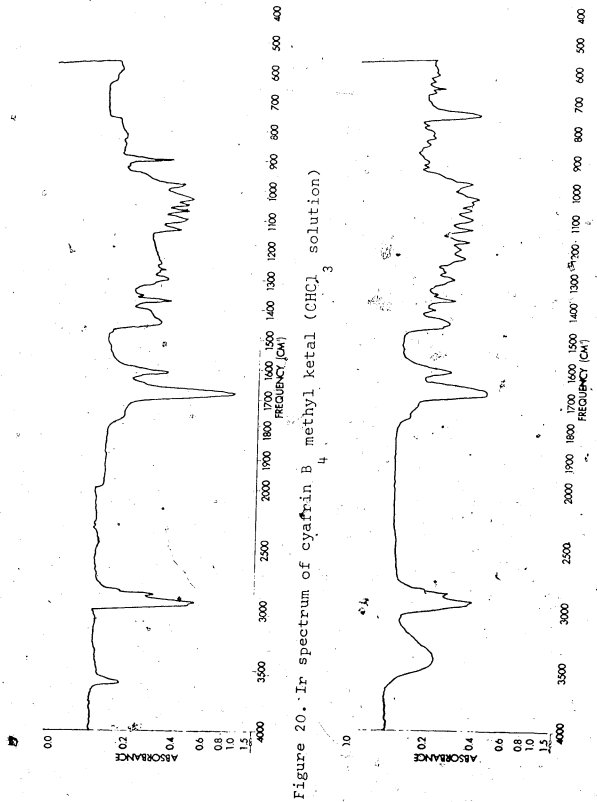


figure 19. Mass spectrum of cyafrin B methyl ketal

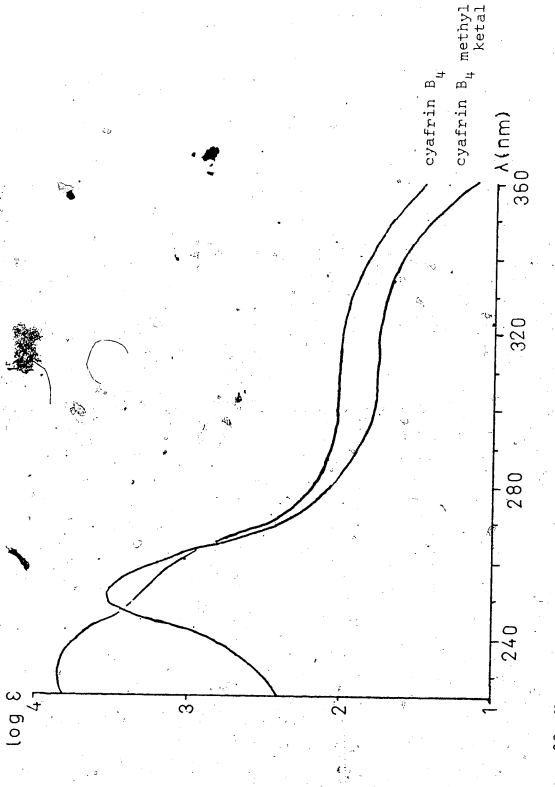
The ir spectrum of cyafrin B methyl ketal in chloroform solution (see fig. 20) shows a strong absorption at 1685 cm indicating the presence of a carbonyl group. Fig. 21 shows the spectrum as recorded on a neat (solid phase) sample of cyafrin B methyl ketal. It is possible that the absorption found at 1685 cm in the solution spectrum of cyafrin B itself (fig. 14) is in fact due mainly to this second carbonyl group and not to the α,β-unsaturated ketone as assumed previously. Interestingly, the solution spectrum of cyafrin B methyl ketal (fig. 20) shows only free hydroxy group absorption (3600 cm ) and ne hydrogen bonded hydroxyl group absorption (3000 cm all, whereas the reverse is true for the solid phase ir spectrum of cyafrin B methyl ketal (fig. 21).

The uv spectrum of cyafrin B methyl ketal in hexane solution proves conclusively that cyafrin B contains a second chromophore besides the  $\alpha,\beta$ -unsaturated ketone common to all cyathins. An absorption is found at 251 nm ( $\epsilon$  ~ 3.500), with a shoulder around 310 nm ( $\epsilon$  ~60). This indicates that a second  $\alpha,\beta$ -unsaturated ketone is present. This absorption must have been partially masked in the uv spectrum of cyafrin B itself. A logarithmic plot of the uv spectrum of cyafrin B reveals a shoulder around 251 nm on the peak at 231 nm (see fig. 22). Thus it has now become clear that cyafrin B contains two  $\alpha,\beta$ -unsaturated ketone functions.



 $_{
m u}$  methyl ketal (neat) Figure 21. Ir spectrum of cyafrin B





and cyafrin  $B_{\mu}$  methyl ketal Figure 22. Uv spectra of cyafrin B

If we assume cyafrin B to possess a cyathinlike structure it is possible to suggest structure <u>21</u>, or alternatively structure <u>22</u> for this metabolite.

The nmr spectrum of cyafrin B methyl ketal (see fig. 23) allows us to choose between 21 and 22. Compare the data for cyafrin B methyl ketal (table X) with the nmr data measured for cyathin A methyl ketal by 7, p.65; 9
Taube (table XI).

The nmr of cyaffin B methyl ketal shows

(table X) signals a, b, c, d, e, q, r, s, t and z which are
all virtually unchanged from their positions in the nmr

spectrum of cyathin A methyl ketal. All other signals of

cyafrin B methyl ketal have shifted to lower field, as was

found in the case of 0,0-diacetylcyafrin B. The two proton

singlet j', found at 62.21 in 0,0-diacetylcyafrin B is

Figure 23. Nmr spectrum of cyafrin B methyl ketal (CDC1 )

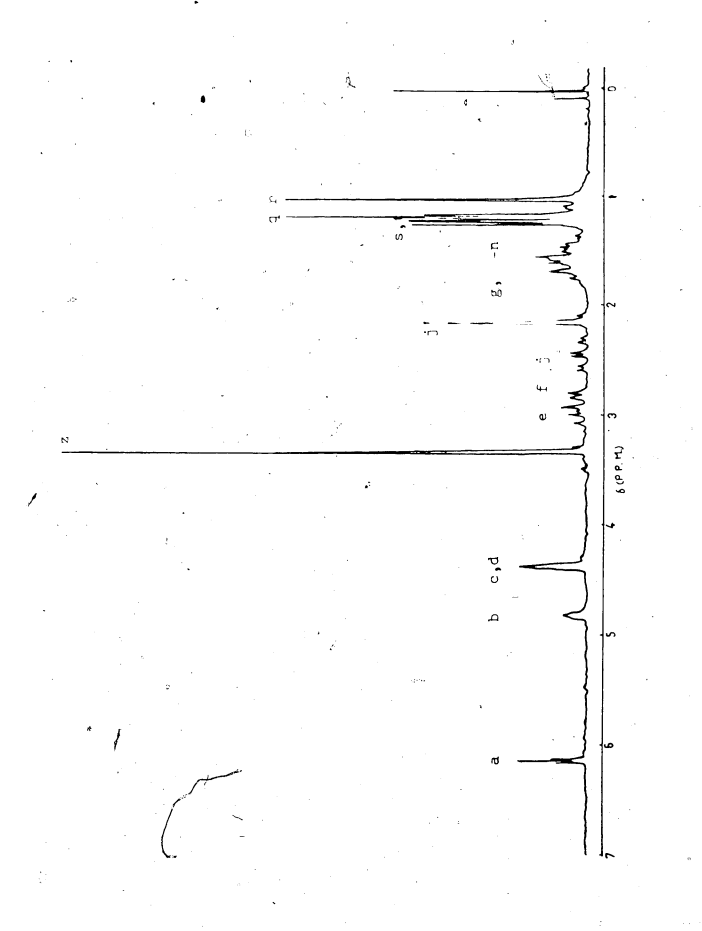


Table X. Nmr data of cyafrin  $B_{\mu}$  methyl ketal in CDCl  $_{\mu}$ 

				•
Coupling constant (Hz)	. S. S.	7,7 12.5,4 12.5,12.5,3.5	· ·	
Multiplicity	ב ב +	ი. ი.ი.ი ი.ი.ი ი.ი.ი ი.ი.ი	ា	מ ס ט
			-	•.
Number of protons	T T Z	<b>പപറ</b>	about 5 3 3	
Chemical shift (6)	6.15 4.82 4.38	3.00 2.89 2.45 2.16	1.8-1.2 1.18 1.01	1.22 1.19 3.34
Signal	а Б С, ф	Ф <del>И</del>	а, к-л д	2 t w

CDC1
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Table

			•				
:	a Assignment	. 13	12	യഗ	2 10 1,7,8,10	(16, 17) (19,	21
	Coupling constant (Hz)	1.5	v. (	6.5,6.5 12,?	12,12,3	ი ი	
tal in CDC1 3	Multiplicity	; t	, ਹ ਹ ਹ	ታ <b>ቴ</b> ታር	ი <b>,</b> ი ი ი ი ი	ם פיטי	ဟ
n A methyl ket 3	Number of protons	H.	2		1 L C	തെതത	സ
Table XI. Nmr data of cyathin A methyl ketal in CDCl	Chemical shift (6)	6.06 4.79	4.34	around 2.42 around 2.2	2.25 1.8-1.2 1.01	0.97 1.03 0.93	3.31
Table XI	Signal	هم <sub>.</sub>	ာ <b>ဝ</b> ိဝ	ب. ئ	j g, k-p q	י ר מ ג	7

Annumbers refer to numbered positions in structure 23 (cyathin A methyl ketal).

present in cyafrin B methyl ketal (table X) at  $\delta$ 2.16. This establishes that signal j' does indeed belong to the compound and is not due to an impurity. The chemical shift of this signal is in accordance with the value expected for the methylene protons adjacent to a carbonyl group in an  $\alpha$  ,  $\beta$  -unsaturated ketone such as those on C-1 in structure 21. Signal d' in the spectrum of 0,0-diacetylcyafrin B (table IX) corresponds to signal f in the spectrum of cyafrin B methyl ketal (table X) and cyathin A methyl ketal (table XI) and may be attributed to the proton on C-5. Its position in structure 21 in relation to the  $\alpha,\beta$ -unsaturated ketone explains why signal d'/f has shifted downfield compared to the values found for cyathin A derivatives. The hydrogens responsible for signals h and i in cyathin A methyl ketal (table XI) have of course been replaced by an oxygen atom in cyafrin B . Also, two protons of the unresolved multiplet around  $\delta$  1.55 in the spectrum of cyathin A methyl ketal (table XI), i.e. those on C-1, now give rise to signal j' in the spectrum of cyafrin B methyl ketal (table X).

Structure  $\underline{22}$  can be eliminated as a possibility for cyafrin B as it would not be expected to show a two proton singlet at  $\delta$  2.21 (signal j') in its nmr spectrum. It should display a signal around  $\delta$  6 due to the vinyl proton on C-2. We are thus led to propose structure 21 for

cyafrin B

It should be noted here that cyafrin B is not identical with the oxidation product obtained previously from cyafrin A (see page 50). This provides additional evidence that in cyafrin A the hydroxyl group is not located on C-2, as was concluded before on the basis of spectral evidence.

## Other metabolites

Throughout this investigation it had been noted that <u>C</u>. <u>africanus</u> produces various other compounds in addition to the four major metabolites discussed thus far. These were sometimes encountered as impurities during isolations of cyafrin A and B and sometimes during mass spectral analysis of bands from prep. tlc which did not contain the major metabolites. Most of these were found only in very small quantities, and usually were not analyzed beyond their molecular formulas (by hrms). Below is a list of the apparent formulas of the minor compounds encountered, arranged roughly in order of increasing R value (in solvent system A, triple elution).

R 0.0-0.2: C H O , C H

C H O . 26 40 8

R 0.2-0.4: C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 , C H 0 . 20 30 5 20 32 4

R 0.4-0.6: C H O , C H

C H O , C H O , C H O . 20 28 5 20 30 4 22 32 4

R 0.6-0.8: C H O , C H O . f 22 30 5 22 32 5

R 0.8-1.0: C H O , C H O , C H O. f 21 30 3 21 32 3 30 46

This compound was encountered increasingly in samples of cyafrin B. If it is isolated in pure form it will not fit conveniently into our present nomenclature system.

Of all these compounds only the ones between R 0.8 and 1.0 have been examined in any detail.

A sample of C H 0 (containing some 21 30 3 C H 0 according to ms) was analyzed using nmr and ir; it 21 32 3 was found that the spectral data were identical with those of allocyathin B methyl ketal, as prepared by Taube , possibly contaminated with some cyathin A methyl ketal. It was then realized that methanol had been used to aid dissolution of the crude cyafrin from which this compound had been isolated; apparently this treatment had been sufficient to convert some of the cyathins present into their

methyl ketal derivatives. In later/separations, when use of methanol was avoided, these compounds were no longer detected.

The other compound that was studied further is the one with apparent molecular formula C H O. Its mass spectrum is shown in fig. 24. Exact mass determination of the fragment of m/e 229 gave as atomic composition C The ir spectrum of this compound showed, besides C-H and C-C absorption bands, a strong absorption band at 1720 cm which indicates the presence of an unconjugated ketong )function. Uv spectroscopy showed an absorption maximum at 245 nm, but this was of such low intensity ( $\epsilon$  ~1500) that it may well have been caused by an impurity. The total ... amount of this compound present in chude cyafrin was too small to allow a more detailed study at the time. Recently , it has been shown . that this compound, the major metabolite to be found in the mycelium of C. africanus, is glochidonol (structure 24), molecular formula C 30 48 2

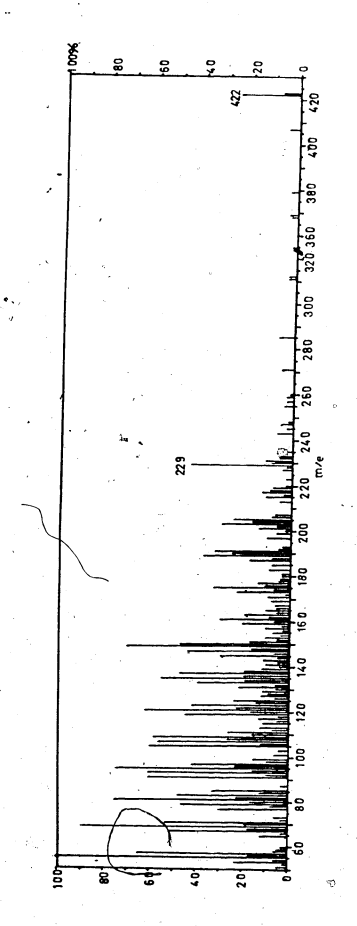


Figure 24. Mass spectrum of glochidonol

## Metabolites of Sphaerobulos stellatus

During the screening of the metabolites of a number of species of bird's nest fungi it was found that a crude ethyl acetate extract of <u>S</u>. stellatus showed some antibacterial activity. It was decided to investigate these metabolites as well.

S. stellatus was cultured under the same conditions as described for C. africanus. Crude ethyl acetate extracts were obtained as described for C. africanus. In addition, the extracted culture was made basic to pH 10 with dilute sodium hydroxide and extracted with methylene chloride to obtain any basic components.

The total amount of crude material isolated from <u>S. stellatus</u> per liter of medium extracted was rather low (only about 30 to 50 mg per liter) compared to the 75 to 150 mg per liter we obtained from <u>C. africanus</u>.

The crude basic extracts were examined by tlc (solvent system C) and found to contain mainly highly polar material (R 0.0) and one small spot at R 0.1. A f very small quantity of this latter material was isolated by prep. tlc (solvent system C). Mass spectroscopy indicated a molecular weight of 168, apparent molecular formula C H N . It was not possible to obtain a good ir spectrum 11/8 2

on this small amount of material. Surprisingly for a compound that appears to be highly unsaturated, we could not detect my significant absorption in the uv spectrum over 220 nm. Later cultures of the fungus did not produce this compound, and it was not further investigated.

Preliminary investigation by tlc (solvent system C) of the crude ethyl acetate extracts revealed a complex mixture, most of it with a low R value, but also f containing some less polar material. Partial separation of this mixture was achieved by means of a distribution between equal volumes of methanol (containing 5% water) and Skellysolve B (a low boiling mixture of hydrocarbons). This resulted in a large "polar fraction" and a much smaller "non-polar fraction".

The "non-polar fraction" was examined by mass spectroscopy and uv. It was found to contain mainly dioctyl-phthalate, a commonly used plasticizer which is readily recognizable from ver prominent peak in the mass spectrum at m/e 149 and other peaks at m/e 167 and 279. No other compounds could be isolated in amounts sufficient for investigation.

The much larger "polar fraction" was subjected to tlc examination using various solvent systems, with a view to separating it by prep. tlc. However, none of the

systems used appeared to be capable of separating the constituents. Usually a streak of material was found on tlc rather than separated spots.

An attempt was made to achieve some separation of this mixture by column chromatography, using silicic acid as adsorbent, and eluting with chloroform - methanol mixtures containing an increasing percentage of methanol (up to 20%). Numerous fractions were collected, and those containing visible amounts of material after evaporation of the solvent were subjected to mass spectral analysis. Peaks were found at m/e 386, 368, 260, 170 and 103. However, during an attempt to obtain the atomic composition of these peaks only the peak at m/e 103 was reproduced. Its molecular composition was found to be C H O . The total amount of material isolated 4 7 3 in this way was very small; the bulk of the material could only be recovered from the column by eluting with pure methanol.

It thus appeared that most of the material was highly polar. It was considered that acetylation of this polar mixture would result in reduction of its polarity and thus perhaps facilitate separation. Also this would facilitate the use of gas chromatography (gc) as an analytical and preparative method.

Acetylation of the mixture was achieved in the

usual way, using a large excess of pyridine - acetic anhydride overnight at room temperature. The reagents were removed by adding toluene and distilling the mixed solvents under reduced pressure, repeating this several times until the residue appeared to be free of the reagents.

The acetylated mixture was used for trial separations using gas chromatography (gc). The best separation was obtained using an OV-225 column (3% on chromosorb W) of 7' x ¼" and temperature programming (100 -220 C at 2 C per minute). In this way as many as thirteen peaks could be detected (fid detector) in the mixture, one of which (at a short retention time) was by far the largest.

Preparative gc was first investigated as a method of isolating this compound. However, it was found that material collected in this way, when re-examined by analytical gc, no longer contained this component; apparently it decomposes during gas chromatography.

Prep. tlc was next investigated, separation: being monitored by isolation of separated bands and then analysis of the mixtures obtained from each band by gc. Of several solvent systems tried none gave a clean separation.

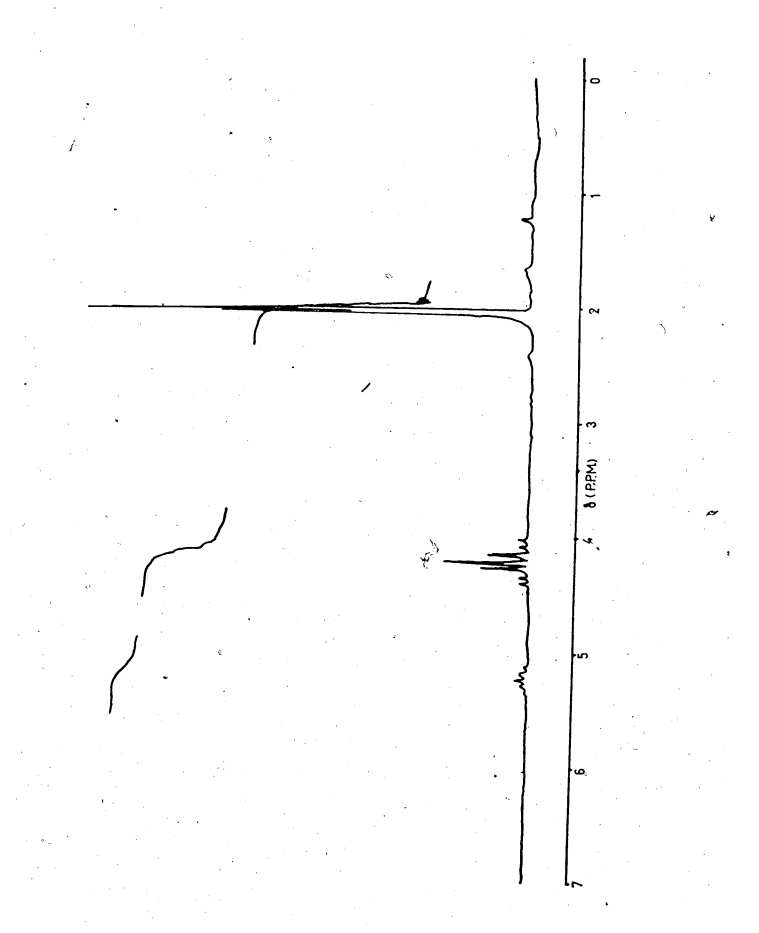
Finally the major component was isolated by column chromatography, using silica gel G as adsorbent and

eluting it with pure chloroform containing approximately 0.75% ethanol as stabilizer. Under these conditions it was the first compound eluted.

The mass spectrum of the compound shows peaks at m/e 158, 145, 116, 115, 103, 87 and 43. Aside from the base peak at m/e 43, the strongest peak is the one at m/e 103 which has an atomic composition of C H O. The nmr spectrum (see fig. 25) is very simple; a double peak at  $\delta 2.05$ , integrated at nine protons, a four proton triplet at  $\delta 4.20$  and a one proton quintet at  $\delta 5.22$ . The ir spectrum showed absorption peaks at 955, 1020, 1060, 1090, 1265, 1370, 1420 and  $\frac{1}{1000}$  No uv absorption maxima were found above 210 nm.

The spectra proved to be identical with those of triacetylglycerol. Glycerol is one of the components of the Brodie medium used for growing S. stellatus. It thus appeared that the most abundant compound in the crude extracts was not a metabolite produced by the fungus. Since minor compounds were amenable to separation and since we had no indication of the presence of cyathin-like compounds, the extracts were not examined further.

Figure 28. Nmr spectrum of triacetylglycerol (CDC1)



### IV. DETAILED EXPERIMENTAL

#### General remarks

The experimental data given in this chapter are in many cases examples of experiments which were carried out several times. The amounts used and the yields varied.

Spectral data presented in chapter III are not repeated, but reference is made to the page where they are recorded.

## Growth of fungi

The procedures for growing the fungi that
were investigated are described under General Experimental.
The Brodie medium used contains per liter: glycerine, 6 ml;
peptone, 0.2 g; d,l-asparagine, 0.2 g; yeast extract, 2.0 g;
MgSO (anhydrous), 0.24 g; Ca(NO) .4H 0, 0.5 g; KH PO, 0.5

4 g; Fe (SO), trace; maltose, 5.0 g; dextrose, 2.0 g. For

2 4 3
liquid cultures this was made up to one liter with distilled
water. For solid media 15 g bacto-agar was added, and distilled water to make up one liter.

The procedure used for preparing crude extracts is also given under General Experimental.

## Isolation of crude metabolites

Crude cyafrin was freed from highly polar material by the ether - water partition procedure described previously. The resulting partially purified cyafrin was dissolved in a small volume of methylene chloride and chromatographed on prep. tlc plates of 100 cm length, applying 0.5 g cyafrin to each plate. The plates were developed with solventsystem A (triple elution). Three bands, at R 0.2, f 0.4 and 0.5 were removed from the plates and eluted to yield, after evaporation of the solvent on a rotary evaporator, crude cyafrin A, crude cyafrin B and crude cyathin A / allocyathin B mixture respectively.

# Acetylation and purification of the crude cyathin $A_3$ / allocyathin $B_3$ mixture

Crude cyathin A / allocyathin B (50 mg) was 3 dissolved in 2 ml of pyridine. Acetic anhydride (1 ml) was added and the mixture was allowed to stand overnight at room temperature. Toluene (15 ml) was added and the mixed solvents were removed by evaporation under reduced pressure using a rotary evaporator. This toluene treatment was repeated three times. The residue was taken up in methylene chloride and chromatographed on one 20 x 20 cm prep. tlc plate (solvent system A). A band at R 0.7 was removed from the plate and eluted to yield, after evaporation of the solvent, 15 mg of

a 3: 4 mixture of 0,0-diacetylcyathin A and 0,0-diacetyl3
allocyathin B. The mass spectrum of this mixture is shown
3
in fig. 1. The nmr spectrum is given in fig. 2.

# Purification of cyafrin $A_4$ and cyafrin $B_{\mu}$

Crude cyafrin A (200 mg, isolated from several growths of fungus) was dissolved in a small amount of acetone and applied to two 20 x 20 cm prep, tlc plates. These were developed with solvent system D. The band at R 0.6 was fremoved from the plates and eluted with acetone. After evaporation of the solvent 140 mg of purified cyafrin A was obtained. The mass spectrum of this compound is shown in fig. 3. Ir spectra are shown in fig. 5 and 6. The nmr spectrum is shown in fig. 4.

Crude cyafrin B (300 mg, isolated from several growths) was dissolved in a small amount of acetone and chromatographed on three 20 x 20 cm. prep. tlc plates with solvent system C, quadruple elution. The band at R 0.4, after elution with acetone and evaporation of the solvent, yielded 94 mg purified cyafrin B. The mass spectrum of this compound is shown in fig. 13. Ir spectra are shown in fig. 14 and 15. The uv spectrum is shown in fig. 22.

## Acetylation of cyafrin $A_{ij}$ and cyafrin $B_{ij}$

Cyafrin B (30 mg) was acetylated in the same manner. After removal of the excess reagents prep. tlc (solvent system A) yielded 0,0-diacetylcyafrin B (21 mg). Mass spectral analysis of this material showed it to be impure. Further purification was achieved by prep. tlc (solvent system B, double elution) on a 5 x 20 cm plate. A band at R 0.5 f yielded pure 0,0-diacetylcyafrin B (8 mg). The mass spectrum of this compound is shown in fig. 16. The ir spectrum is shown in fig. 17. Nmr data are given in fig. 18 and table IX.

# Formation of acetonide of cyafrin Au

thoxypropane ( to which a few drops of acetone had been added to assist d. on. A few small crystals of p-to-

luenesulphonic acid were added. After 30 minutes the reaction mixture was applied directly to a 5 x 20 cm prep. tlc plate, which was developed with solvent system A. The band at R 0.5 f yielded pure 0,0-isopropylidenecyafrin A (9 mg). The mass spectrum of this compound is shown in fig. 9. Nmr data are given in fig. 10 and tables IV and VI.

# "Jones" oxidation of 0,0-isopropylidenecyafrin $A_{ll}$

0,0-isopropylidenecyafrin A (5 mg) was dissolved in acetone (0.2 ml). "Jones reagent" (8N Cro in dilute sulphuric acid, 15 microliter) was added. The orange colour of the reagent changed to green. A small drop of 2-propanol was added to consume excess reagent and a small amount of Na Co was added to neutralize the solution. The solids were filtered off and the reaction mixture was applied to a 5 x 20 cm prep. tlc plate. After development with solvent system A, the band at R 0.75 yielded dehydrocyafrin A acetonide (approx. 1 mg). For analytical data on this compound see page 50.

# Attempted dehydration of 0,0-isopropylidenecyafrin A4

0,0-isopropylidenecyafrin A (5 mg) and collidine (0.3 ml) were dissolved in a small amount of methylene chloride. Alumina (Woelm neutral alumina - grade 1, 0.5 g) was added, and the methylene chloride removed by evaporation.

The resulting dry powder was heated at 160°C for 30 minutes, then extracted first with 10 ml methylene chloride and then with 10 ml of methanol. The combined extracts were concentrated by evaporation. The examination of the residue (solvent system A) showed no spots with R greater than 0.0. No starting material could be detected. This material was not further characterized.

## Formation of 1-chlorocyathin A3 acetonide

0,0-isopropylidenecyafrin A (7 mg) was dissolved in dry acetone (0.5 ml). Pyridine (20 microliter) was added and the solution was cooled to 0 C in an ice-water bath, then 10 microliter of freshly distilled thionyl chloride was added. After 50 minutes the reaction mixture was applied directly to a 5 x 20 cm prep. tlc plate, which was developed with solvent system A. The band at R 0.8 yielded what is believed to be 1-chlorocyathin A acetonide (2 mg). The mass spectrum of this compound is shown in fig. 11. The nmr spectrum of this compound is shown in fig. 12.

# Formation of methyl ketal of cyafrin $B_{4}$

Cyafrin B (90 mg, impure) was dissolved in 4.5 ml of a 2.5 N solution of HCl in methanol. After 4 hours the reaction mixture was applied to a 20 x 20 cm prep. tlc plate. After development with solvent system A (double elu-

tion) a band at R 0.4 gave cyafrin B methyl ketal (45 mg). f

The mass spectrum of this compound is shown in fig. 19. The ir spectra are shown in fig. 20 and 21. The uv spectrum is shown in fig. 22. Nmr data are given in fig. 23 and table X.

## Isolation of triacetylglycerol

Some of the methanol soluble fraction of the ethyl acetate extract from S. stellatus was acetylated as described for cyathin A / allocyathin B. About 100 mg of 3 this acetylated crude was chromatographed on a column of silica gel G (E. Merck), 1 cm diameter, 10 cm high. The column was eluted with chloroform (containing approx. 0.75% ethanol as stabilizer). The fractions which were collected were analyzed by glc, under conditions described on page 97. The first five fractions yielded, after evaporation of the solvent, glycerol triacetate (33 mg). The nmr spectrum of this compound is shown in fig. 28.

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