

P A R E - III

P A R T . III

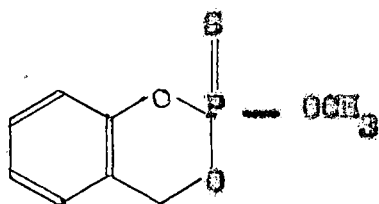
STUDIES ON SOME SALIGENIN CYCLIC PHOSPHORAMIDOTHIONATES HAVING  
FUNGICIDAL ACTIVITIES.

[ Synthesis, fungicidal activity, insecticidal activity, acute oral toxicity, phytotoxicity, anticholinesterase activity, and chemical hydrolysis of some 6-nitro-saligenin cyclic phosphoramidothionates. ]

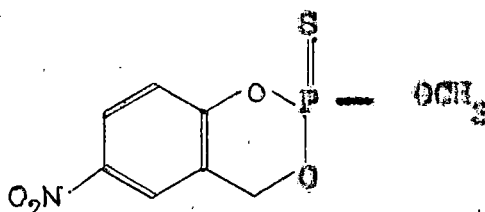
### PART - III

#### AIMS AND OBJECTS OF THE PRESENT INVESTIGATION.

As stated previously (in Part - II) the cyclic organophosphorus esters of saligenin were discovered as the biologically active metabolites of tri-ortho-tolyl phosphates; many related compounds have been synthesized to study their chemical, biochemical and biological properties. Salithion (2-methoxy - 4H - 1,3,2 - benzodioxaphosphorin - 2 - sulphide) is now commercialised as an insecticide. The introduction of any type of substituent at any position of the benzene ring decreases the insecticidal activity <sup>(1)</sup>. It has been reported <sup>(1)</sup> that the BD-8 (2 - methoxy - 6 - nitro - 4H - 1,3,2 - benzodioxaphosphorin - 2 - sulphide) is obtained as a paste in the reaction of O - methyl-dichloridophosphorothionate with 5 - nitro saligenin after purification through silicic acid column chromatography; and, this methoxy compound (BD-8) <sup>(1)</sup> has about sixty times less insecticidal activity compared to salithion.



Salithion

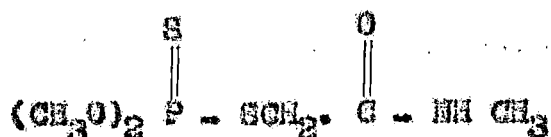


BD-8 (m.p. --- 84°C)

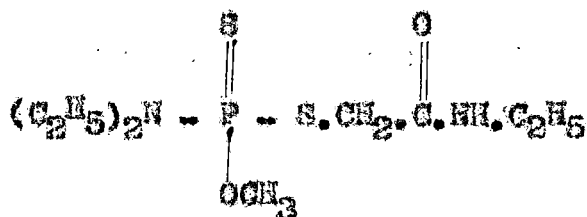
However, it has been observed <sup>(2)</sup> in this laboratory that the methoxy compound (BD-8) is a solid (m.p. 84°C), and has about 1.5 to 2 times greater oral insecticidal activity to cockroaches, Periplaneta americana (Linn) compared to salithion; and also its (BD - 8's) contact

insecticidal activity against grasshoppers, Oxya nitidula is comparable with that of salithion<sup>(3)</sup>. Our investigations on 6-nitro saligenin cyclic phosphorus compounds have started with the finding of insecticidal activity in the compound BA-8.

Introduction of an amide group in place of an alkyl ester group often affords organophosphorus esters with fungicidal, herbicidal, and some-times insecticidal activity<sup>(4)</sup>. Phosphoramidates having a bulky alkyl group as iso-propyl, secondary-butyl, and cyclohexyl on the amido nitrogen atom show high herbicidal activity, whereas N-ethyl, N-methyl or N-monosubstituted phosphoramidate are suitable as insecticide. Although the insecticide dimethoate [dimethyl S-(N-methylcarbamoyl methyl) phosphorothiothionate]<sup>7</sup> has no fungicidal activity, its dialkylphosphoramidate analogs such as compound - I,<sup>(5)</sup> show some fungicidal as well as acaricidal activity.



Dimethoate



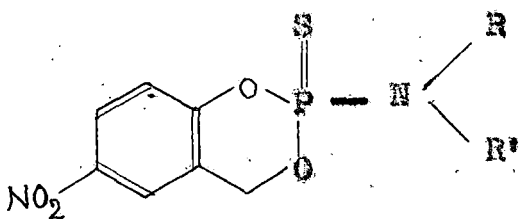
Compound - I

The S-alkylcarbamoylmethyl group is not required for the fungicidal effect, and some S-phenylphosphoramidethiothionates are more active than compound - I. Phosbutyl (ethyl S-phenyl N-butylphosphoramidethiothionate)<sup>(6)</sup> is a practically useful fungicide in this series. An amido analog of Kitazin, S-benzyl ethyl N-isopropylphosphamido-

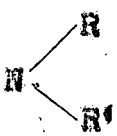
thiolate) <sup>(7)</sup> was patented as a fungicide .

There are several other examples in literature from which it can be found that some phosphoramidothionates, phosphoramidothiothionates, phosphoramides or phosphonamides in which the phosphorus atom is attached directly to the nitrogen atom of an amine or a heterocyclic compound such as phthalimide, imidazole or triazole have very good fungicidal activity <sup>(4,6,8,9)</sup> .

These observations prompted us to undertake a systematic work on some 6 - Nitro saligenin cyclic alkylamidophosphorothionates having the general structure (A),



(A)

where,  is cyclo-hexylamido, morphilino, dimethylamido, diethylamido, pyrrolidino, piperidino, isopropylamido, nonylamido or heptadecylamido. The work embodied in this dissertation is related to the investigation of the above mentioned compounds with reference to their chemical, fungicidal, insecticidal, toxicological, anticholinesterase, phytotoxic and hydrolytic properties besides structural elucidations by spectroscopic methods.

R E F E R E N C E S.

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A : MATERIALS AND METHODS

## A. MATERIALS AND METHODS

### 1. PURIFICATION OF SOLVENTS AND OTHER CHEMICALS:

Throughout the preparative and other part of the work, the organic solvents and other chemicals used were purified and dried according to Vogel <sup>(1)</sup>. All other chemicals and solvents occasionally used during the work were of standard commercial products of high quality (EM, BDH, SM and/or Fluka quality).

### 2. SPECTROSCOPIC METHODS:

Infrared spectra were obtained with Beckmann IR-20 spectrophotometer in nujol mull. PMR spectra were recorded on Varian Model A - 60, EM - 390 and FT - 80A instruments in  $CDCl_3$  solvent using TMS as an internal reference. Mass spectra were taken on Varian Model EM - 600 and MS - 30 instruments.

### 3. PREPARATION OF THIOPHOSPHORYL CHLORIDE ( $PSCl_3$ ):

Thiophosphoryl chloride was prepared according to Moeller <sup>(2)</sup>  
et al

### 4. PREPARATION OF 2 - NITRO - SALIGENIN ( 2 - HYDROXY - 5 - NITRO - BENZYL ALCOHOL ):

The 2 - hydroxy - 5 - nitrobenzyl alcohol as one of the starting materials for the synthesis of nitro-saligenin cyclic alkylamido phosphorothionates was prepared in the following manner:

Preparation of the alcohol was done in two stages:

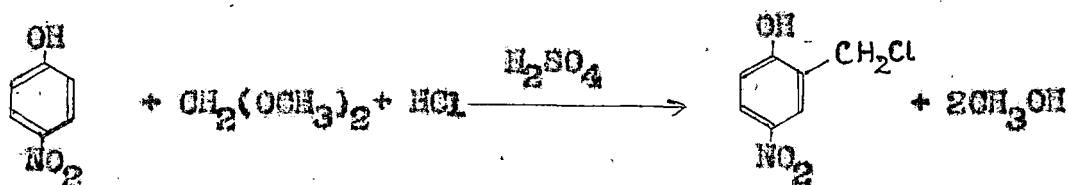
(1) Preparation of the 2 - hydroxy - 5 nitro benzyl chloride,

and

(ii) the hydrolysis of the said chloride.

4.(1) PREPARATION OF 2-HYDROXY - 5 - NITROBENZYL CHLORIDE:

The 2 - hydroxy - 5 - nitrobenzyl chloride was prepared according to the method described in organic synthesis <sup>(3)</sup>. The reaction involved is :



The p - nitrophenol melting at above 112°C (Reidel, m.p. 114°C) was used for the preparation. The other ingredient methylal was synthesized afresh for every batch of preparation as follows:

760 ml. methanol was added to 400 gm anhydrous calcium chloride in a 3 lt. round bottomed flask equipped with a reflux condenser, 10.2 ml. conc. hydrochloric acid was added; then with cooling and constant stirring, 400 gm of 37 - 40% formaldehyde was slowly dropped in through a dropping funnel. It took about 2 hours for complete addition of formaldehyde (the reaction was strongly exothermic). When all the formaldehyde had been added, the mixture was heated for a few minute until the liquid boiled vigorously. The methylal came up

quickly on the upper layer, and after an overnight standing was fractionally distilled. The 42 - 45°C fraction was collected and stored in a tightly stoppered bottle in cold (freezer) before it was used.

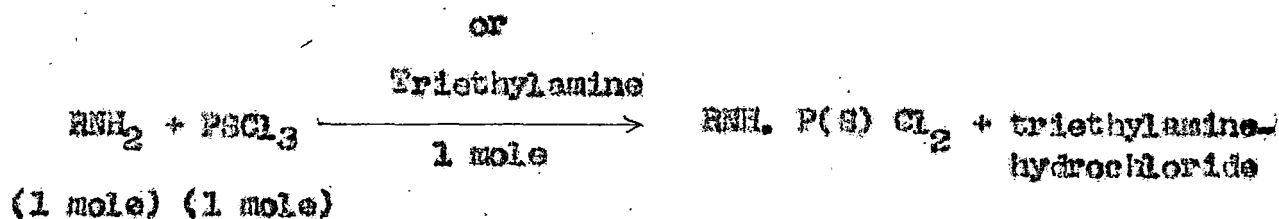
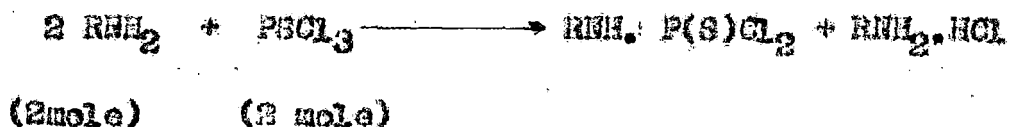
In a one-litre, three-necked round bottomed flask equipped with a mechanical stirrer, a short reflux condenser, and a bent glass tube reaching sufficiently below, were placed 50 gm (0.26 mole) p - nitrophenol, 650 ml conc. HCl, 5 ml conc. H<sub>2</sub>SO<sub>4</sub> and 76 gm (1 mole) methylal. The reaction mixture was stirred while the temperature was maintained at and around 70 ± 2°C for 4 - 5 hrs. by means of a water bath. During this time HCl gas was bubbled into the reaction mixture through the bent glass tube. The 2 - hydroxy- 5 - nitrobenzyl chloride began to separate as a solid after about one to one and half hour. At the end of the reaction, the mixture was cooled in a ice - bath for a period of 1-2 hour, more crystals separated. The solid materials, after filtration, were air dried for several hours to remove water; and then washed with benzene. The yield was about 40 - 45 gm. of a white product, m.p. 129 - 130°C.

#### 4. (11) HYDROLYSIS OF 2 - HYDROXY - 5 - NITROBENZYL CHLORIDE.

The said chloride was taken as suspension in water and heated slowly; the crystals would go into solution while the impurities form a deep brown oily layer at the bottom of the container. The mixture was then boiled gently to ensure complete hydrolysis. The solution was quickly filtered while hot. The light yellow crystals of 2 - hydroxy - 5 - nitrobenzyl alcohol separated as the filtrate got cooled. The product (m.p. 122 - 126°C) thus obtained was recrystallised from hot water; the

crystals were filtered, washed with cold dioxan; benzene ( 1:9) mixture, and dried in vacuum.

5. PREPARATION OF PHOSPHORAMIDOTHIOIC DICHLORIDES:



One mole  $\text{PSCl}_3$  and two moles amine (or one mole amine and one mole triethylamine) were allowed to react at  $+5^\circ$  to  $-5^\circ\text{C}$  in benzene/chloroform as solvent. The amine solution was added dropwise very slowly with constant vigorous stirring. After an additional stirring period, the solid particles were filtered off the reaction mixture and the liquid phase was washed repeatedly with benzene/chloroform, subsequently with 2% ~~wt~~ cold hydrochloric acid, then with cold saturated solution of sodium chloride. The benzene/chloroform phase was then dried with anhydrous sodium sulphate and filtered; evaporation in Vacuo gave the desired phosphoramidothioic dichloride; this dichloride was used as such for the subsequent reaction.

The different phosphoramidothioic dichlorides were prepared as follows:

5.(1) CYCLOHEXYLAMIDOPHOSPHORODICHLORIDOTHIONATE:

Thiophosphoryl chloride (16.9 gm; 0.1 mole) in 100 ml benzene was allowed to react with cyclohexylamine (19.8 gm; 0.2 mole) in 20 ml benzene at  $-20^{\circ}$  to  $5^{\circ}\text{C}$ ; the amine solution was added dropwise very slowly with constant vigorous stirring. After an additional stirring period of two hours, the cyclohexylamine hydrochloride was filtered off, and the solution was washed with 2% cold hydrochloric acid saturated with sodium chloride, and then with cold saturated solution of sodium chloride. The benzene phase was then lyophilised with anhydrous sodium sulphate and filtered; evaporation in Vacuo gave the cyclohexylamidophosphorodichloridethionate. 16 gm (70% yield) white crystalline solid (m.p.  $68 - 69^{\circ}\text{C}$ ) was thus obtained.

5.(11) MORPHOLINOPHOSPHORODICHLORIDOTHIONATE:

A solution of morpholine (8.7 gm; 0.1 mole) in 20 ml benzene was added dropwise to a stirred solution of thiophosphoryl chloride (8.45 gm; 0.05 mole) in 50 ml benzene at  $-5^{\circ}$  to  $+5^{\circ}\text{C}$ . The mixture was stirred at  $5^{\circ}\text{C}$  for 3 hr. and at room temperature for 16 hr. The mixture was worked up as in 5.(1). A clear viscous liquid which solidified after standing several days was obtained (7.0 gm; 64% yield,) m.p.  $30 - 31^{\circ}\text{C}$ ;

5.(111) N,N - DIETHYLAMIDOPHOSPHORODICHLORIDOTHIONATE:

Diethylamine was extracted in benzene from its aqueous solution by partition method, and its concentration was determined by volumetric method; before estimation of the amine the benzene phase

was dried with anhydrous sodium sulphate.

7.3 gm (0.1 mole, 50.12 ml. benzene extract solution conc. 0.1456 gm/ml) diethylamine was allowed to react with thiophosphoryl chloride (8.45 gm, 0.05 mole) in 50 ml. benzene at  $-5^{\circ}$  to  $+5^{\circ}$ C and the mixture was worked up as in 5 (1). 8.0 gm (78% yield) liquid possessing camphor - like odour was thus obtained.

5. (iv) N, N - DIMETHYLAMIDOPHOSPHORODICHLORIDOTHIONATE;

N, N - Dimethylamine was extracted in chloroform from its aqueous solution by partition method and its concentration was determined by volumetric method; before estimation of the amine, the chloroform phase was dried with anhydrous sodium sulphate.

Thiophosphoryl chloride (8.45 gm; 0.05 mole) in 50 ml chloroform was allowed to react with 4.5 gm (0.1 mole, 37.8 ml chloroform extract solution, conc. 0.1190 gm/ml) dimethylamine and the mixture was worked up as in 5.(1). 7.5 gm (84% yield) liquid which solidified on standing as crystals (m.p.  $32 - 33^{\circ}$ C) was thus obtained. This compound has camphor-like odour.

5.(v) ISOPROPYLAMIDOPHOSPHORODICHLORIDOTHIONATE;

Thiophosphoryl chloride (8.45 gm; 0.05 mole) in 50 ml chloroform was reacted with 5.9 gm (0.1 mole, 24.8 ml chloroform extract solution, conc. 0.2379 gm/ml) isopropylamine, and the mixture was worked up as in 5.(1). 8.0 gm (83% yield) white crystalline solid (m.p.  $35 - 36^{\circ}$ C) was thus obtained.

5. (vi) PYRROLIDINOPHOSPHORODICHLORIDOTHIONATE:

Thiophosphoryl chloride (8.45 gm, 0.05 mole) in 50 ml chloroform was reacted with 7.1 gm (0.1 mole) pyrrolidine in 20 ml chloroform, and the mixture was worked up as in 5.(i). 8.5 gm product was obtained

5. (vii) PIPERIDINOPHOSPHORODICHLORIDOTHIONATE:

Thiophosphoryl chloride (8.45 gm, 0.05 mole) in 50 ml chloroform was allowed to react with 8.5 gm (0.1 mole) piperidine in 20 ml chloroform. 9.0 gm product was obtained.

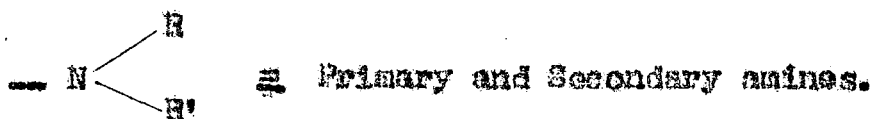
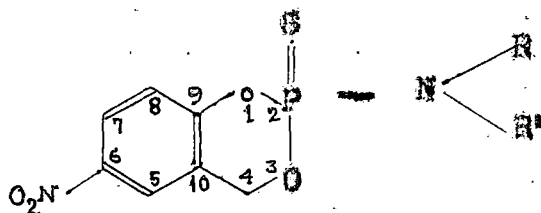
5. (viii) NONYLAMIDOPHOSPHORODICHLORIDOTHIONATE:

This compound was prepared by taking  $\text{PSCl}_3$  (4.1 gm), triethylamine (2.45 gm, 0.0242 mole), and n - nonylamine (3.47 gm, 0.0242 mole) in 100 ml. chloroform; the nonylamine in 30 ml. Chloroform was added dropwise; after additional stirring for 3 hrs. the reaction mixture was washed twice with cold saturated sodium chloride solution. The chloroform phase was then dried with anhydrous sodium sulphate and filtered; evaporation in Vacuo gave 6.2 gm liquid nonylamidophosphorodichloridothionate.

5. (ix) HEPTADECYLAMIDOPHOSPHORODICHLORIDOTHIONATE:

Thiophosphoryl chloride (1.32 gm, 0.0078 mole), triethylamine (0.79 gm, 0.0078 mole), and n - heptadecylamine (1.994 gm, 0.0078 mole) were used for the preparation of this compound; the heptadecylamine in 30 ml chloroform was added dropwise; 1.8 gm product was obtained.

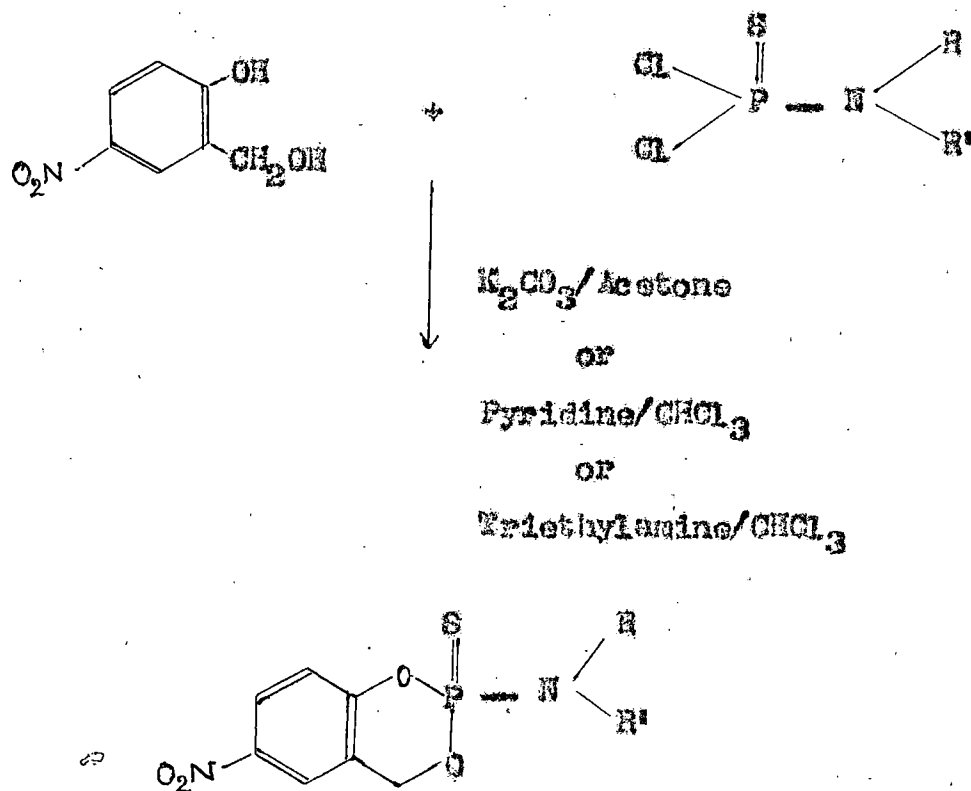
6. PREPARATION OF 2 - ALKYLAMIDO - 6 - NITRO - 4H - 1,3,2 - BENZODIOXA-  
PHOSPHORIN - 2 - SULPHIDE:



6. (a) GENERAL PROCEDURE:

2-alkylamido - 6 - nitro - 4H - 1,3,2 - benzodioxaphosphorin - 2 - sulphides were prepared by adding a solution of 2 - hydroxy - 5 - nitrobenzyl alcohol (5 - nitro saligenin, 1 mole) in dry acetone to 1 mole of alkylamidophosphorodichloridothionate with cooling in an ice bath. The anhydrous potassium carbonate (2 mole) was then added by instalments, with constant stirring. In some cases, good result was found by using 2 moles of pyridine or 2 moles of triethylamine in chloroform solution instead of anhydrous potassium carbonate. The temperature of the reaction mixture was strictly maintained below 5°C during the addition of potassium carbonate/pyridine/triethylamine. The condensation was accomplished by stirring at the temperature 5 - 27°C for an additional time of 12 - 16 hrs. The solid particles were filtered out of the reaction mixture and the solvent was removed under reduced pressure at room temperature. In some cases the crude product was directly re-

crystallised from methanol to give the pure compound; while in others an additional chloroform extraction was necessary prior to the recrystallisation. In the latter case, the crude product was extracted with chloroform and washed with 1% dil HCl (ice - cooled) and with cold water, repeatedly (three times). This was then dried with anhydrous sodium sulphate and the chloroform was subsequently removed under reduced pressure. The pure compounds were then obtained by recrystallisation from methanol.

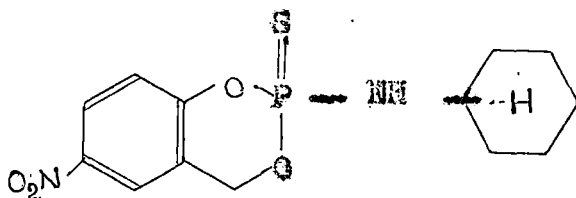


The different alkylamidophosphorothionates were prepared as follows:

6. (b) PREPARATION AND PROPERTIES OF SOME INDIVIDUAL ALKYLAMIDOPHOSPHORODICHLOROTHIONATES:

Melting points are uncorrected, Micro analysis are carried out by Alfred Bernhardt, West Germany.

6. (b) (1) 2-CYCLOHEXYLAMIDO - 6-NITRO - 4H - 1,3,2 - BENZODIOXAPHOSPHORIN-2 - SULFIDE (BD-10):



This compound (BD-10) was prepared by condensation of 5-nitro saligenin (1.69 gm; 0.01 mole) and cyclohexylamidophosphorodichlorodithionate (2.32 gm, 0.01 mole) in presence of  $K_2CO_3$  (2.76 gm; 0.02 mole) in 75 ml acetone as solvent;  $K_2CO_3$  was added by instalments to the stirred solution at  $0^\circ$  to  $5^\circ C$ . After an additional stirring (2 - 3 hrs, at  $0^\circ$  to  $5^\circ C$ , and then 12 - 16 hrs. at room temperature), the solids were filtered off, and the solvent was removed. The crude product was washed with methanol saturated with n - heptane, and then the compound was recrystallised from hot methanol. 2.5 gm (76% yield) compound was obtained. BD-10 is a white crystalline solid, m.p.  $125^\circ C$ , practically insoluble in water, highly soluble in methanol, ethanol and benzene.

Analysis

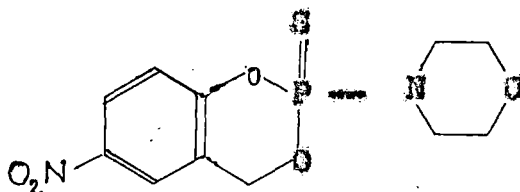
Found : C, 47.54% ; H, 5.19% ; N, 8.52% ;

Calculated for :  $C_{13}H_{17}O_4N_2PS$  : C, 47.56% ; H, 5.18% ; N, 8.53% .

Molecular Weight : 328

A better yield was obtained by using pyridine (chloroform as solvent) instead of  $K_2CO_3$  as dehydrogen chloride agent.

6.(b)(ii). 2 - MORPHOLINO - 6 - NITRO - 4H - 1,3,2 - BENZODIOPHOSPHORIN - 2 - SULPHIDE (BD-11) :



5 - Nitro - Saligenin (1.69 gm, 0.01 mole), morpholinophosphorodichloridethionate (2.2 gm, 0.01 mole), and  $K_2CO_3$  (2.76 gm, 0.02 mole) in 50 ml acetone gave 3 gm (90% yield) crude product; this was then dissolved in chloroform and washed with cold dilute HCl and water; after drying with anhydrous sodium sulphate the solvent was removed under reduced pressure. The solid mass was then recrystallised from hot methanol. 2.0 gm (60% yield) BD-11 was thus obtained; BD-11 is a white crystalline solid, m.p. 149°C, highly soluble in acetone and chloroform, moderately soluble in methanol, ethanol and benzene.

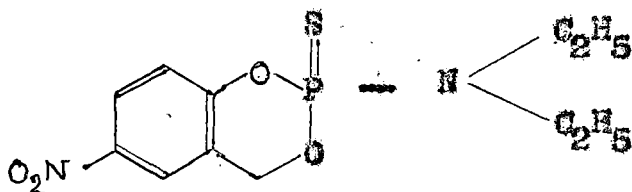
Analysis

Found : C, 41.75%; H, 4.10%; N, 9.24%.

Calculated for :  $C_{11}H_{13}O_5N_2PS$  : C, 41.77%; H, 4.11%; N, 9.27%.

Molecular Weight : 316.

6.(b)(iii). 2 - DIETHYLAMIDO - 6 - NITRO - 4H - 1,3,2 - BENZODIOXAPHOS-  
PHORIN - 2 - SULPHIDE (BD.12):



5 - Nitro - Saligenin (1.69 gm, 0.01 mole), diethylamidophosphorodichloridothionate (2.06 gm, 0.01 mole), and  $K_2CO_3$  (2.76 gm, 0.02 mole) in 50 ml acetone gave BD.12 (2.8 gm, 90% yield) as white crystals (after working up as in BD.10), m.p. 105°C.

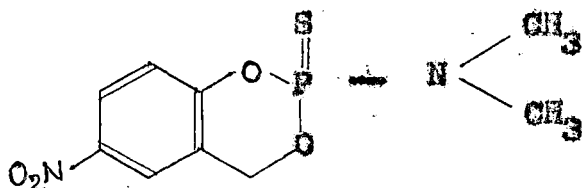
Analysis

Found : C, 43.68%; H, 4.94%; N, 9.25%.

Calculated for :  $C_{11}H_{15}O_4N_2PS$ ; C, 43.70%; H, 4.93%; N, 9.27%

Molecular Weight : 302

6.(b)(iv). 2 - DIMETHYLAMIDO - 6 - NITRO - 4H - 1,3,2 - BENZODIOXA-  
PHOSPHORIN - 2 - SULPHIDE (BD-13):



5 - Nitro - Saligenin (1.69 gm, 0.01 mole); dimethylamidophosphorodichloridothionate (1.78 gm, 0.01 mole),  $K_2CO_3$  (2.76 gm, 0.02 mole) in 50 ml acetone gave BD-13 (2.0 gm, 75% yield) as white crystals (after working up as in BD-10) m.p. 128°C.

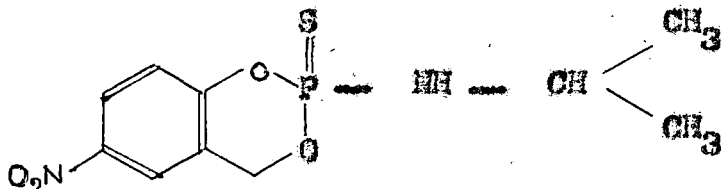
Analysis

Found : C, 39.40%; H, 4.00%; N, 10.19%.

Calculated for :  $C_9H_{11}O_4N_2PS$  : C, 39.41%; H, 4.01%; N, 10.21%.

Molecular Weight : 274.

6.(b)(v). 2 - ISOPROPYLAMIDO - 6 - NITRO - 4H - 1,3,2 - BENZODIOXAPHOS-  
PHORIN - 2 - SULPHIDE (BD-14):



5 - Nitro - Saligenin (1.69 gm, 0.01 mole) isopropylamidophosphorodichloridothionate (1.92 gm, 0.01 mole),  $K_2CO_3$  (2.76 gm, 0.02 mole) in 50 ml acetone gave BD-14 (2.0 gm, 70% yield) as white crystals, m.p.

98°C.

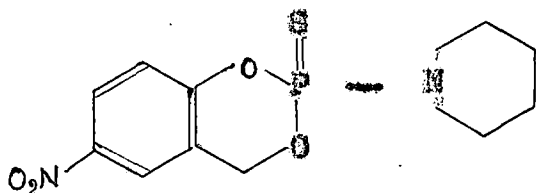
Analysis

Found : C, 41.64%; H, 4.50%; N, 9.70%;

Calculated for :  $C_{10}H_{13}O_4N_2PS$  : C, 41.66%; H, 4.51%; N, 9.72%;

Molecular Weight : 288

6.(b)(vi). 2 - PYRROLIDINO - 6 - NITRO - 4H - 1,3,2 - BENZODIOXAPHOS-  
PHONIN - 2 - SULPHIDE (BD-15):



5 - Nitro - Saligenin (2.70 gm, 0.016 mole), pyrrolidinophosphorodichloridothionate (3.45 gm, 0.016 mole),  $K_2CO_3$  (4.5 gm, 0.032 mole) in 50 ml acetone gave BD-15 (4 gm, 83% yield) as white crystals, m.p. 134°C.

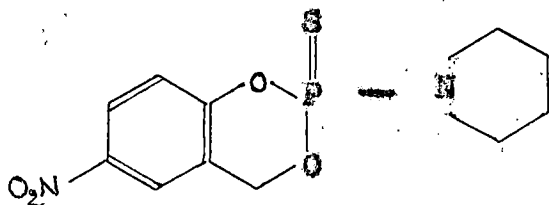
Analysis

Found : C, 43.82%; H, 4.27%; N, 9.23%.

Calculated for :  $C_{11}H_{13}O_4N_2PS$  : C, 43.91%; H, 4.32%; N, 9.32%.

Molecular Weight : 300.6

6.(b) (vii). 2 - PIPERIDINO - 6 - NITRO - 4H - 1,3,2 - BENZODIOPHOSPHONIN - 2 - SULPHIDE (BD -16):



5 - Nitro - Saligenin (3.4 gm, 0.02 mole), piperidinophosphorodichloridothionate (4.36 gm, 0.02 mole),  $K_2CO_3$  (5.5 gm 0.04 mole) in 50 ml acetone gave BD-16 (5.6 gm, 87% yield) as white crystals, m.p.  $130^{\circ}C$ .

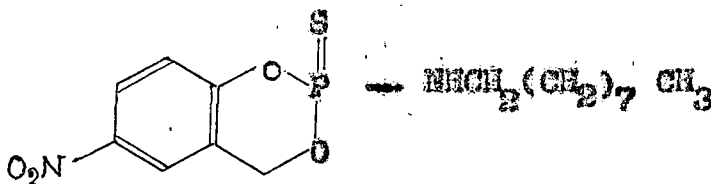
Analysis

Found : C, 45.72%; H, 4.71%; N, 8.82%.

Calculated for ;  $C_{12}H_{15}O_4P_2S$  ; C, 46.71%; H, 4.77%; N, 8.90%.

Molecular Weight : 314.6

6. (b) (viii). 2 - NONYLAMIDO - 6 - NITRO - 4H - 1,3,2 - BENZODIOPHOSPHONIN - 2 - SULPHIDE (BD-17):



5 - Nitro - Saligenin (1.7 gm, 0.0108 mole) nonylamidophosphorodichloridothionate (2.93 gm, 0.0108 mole),  $K_2CO_3$  (2.9 gm, 0.0216

mole) in 50 ml acetone gave BD-17 (3.5 gm, 87% yield) as crystals, m.p. 64°C.

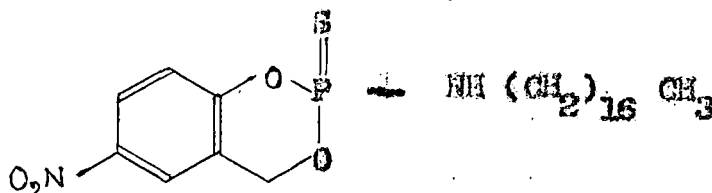
Analysis

Found : C, 51.58%; H, 6.69%; N, 7.50%.

Calculated for :  $C_{16}H_{26}O_4N_2PS$  : C, 51.61%; H, 6.72%; N, 7.53%.

Molecular Weight : 372.

6.(b)(ix). 2 - HEPTADECYLAMIDO - 6 - NITRO - 4H - 1,3,2 - BENZODIOXA-  
PHOSPHORIN - 2 - SULPHIDE (BD-18):



5 - Nitro - Saligenin (0.78 gm, 0.0046 mole), heptadecylamidophosphorodichloridethionate (1.8 gm, 0.0046 mole), triethylamine (0.98 gm) in 50 ml chloroform gave BD-18 (1.82 gm) as crystals, m.p. 70°C.

Analysis

Found : C, 59.46 ; H, 8.41 ; N, 5.76.

Calculated for :  $C_{24}H_{41}O_4N_2PS$  : C, 59.50 ; H, 8.47 ; N, 5.78

Molecular Weight : 484.

7. FUNGICIDAL ACTIVITY:

7. (a) TEST ORGANISM:

Pyricularia Oryzae Cav. - causal fungus of blast disease of rice.

7. (b) CULTURE MEDIUM :

Solid medium : Malt extract agar.

20 gm malt (Difco) extract was boiled in water till dissolved. 20 gm agar agar (Kobe Japan) was added and boiled until agar was well dissolved. 0.05 gm chloramphenicol was suspended in 10 ml of 95% alcohol and added to the medium as antibacterial agent. The volume of the medium was then made upto 1 litre by adding water. pH of the medium was adjusted to 6.5 by adding NaOH solution. Medium was sterilized at 15 p.s.i. for 20 minutes.

7. (c) GROWTH INHIBITION METHOD:

Growth inhibition studies were made by using poison food technique (4). Acetone solution of suitable quantity of the compounds in sterile water containing 0.01 percent Triton - X was incorporated into melted malt agar so as to get the desired concentration of the compounds in the media. The test medium was poured into the sterile petridishes and after solidification the 7 mm 8 days old culture disc was placed aseptically at the centre of the petridish. Three replications on each test with appropriate control under same conditions were maintained. These petridishes were incubated at  $30 \pm 1^{\circ}\text{C}$  in dark. Linear growth of the fungal disc were measured after 24, 48, 72 and 96 hours interval

Percent inhibition over control was calculated following the equation  
(5)  
given by Vincent .

8. INSECTICIDAL ACTIVITY ON GRASSHOPPERS:

Insecticidal tests were also performed on the Sporadic Grasshoppers, Oxya nitidula by topical application of acetone solution of the compounds according to Eto et al (6). The grasshoppers weighing about 0.15 to 0.25 gm, were collected in the month of October - November, 1981 from one particular location of the North Bengal University Campus. In the field they were never exposed to any Organophosphorus Insecticides. For preliminary experiment, acetone solution of the compound was topically applied to ten insects in each pot, and after 24 hours the mortality was determined.

To determine the more precise  $LC_{100}$  value of each compound, one insect of known weight in each pot was treated, by topical application, with acetone solution of test chemicals on mouth and thoracic region (Ventral side) of the insect. Each test was triplicated. For all the alkylamidophosphorothionates the concentration was increased by  $1 \mu\text{g/insect}$ , but for the salithion (standard) the concentration was increased by  $0.1 \mu\text{g/insect}$ .

9. ACUTE ORAL TOXICITY TESTS ON RATS:

Oral toxicity testing was conducted on 6 - 12 months old male white albino rats, weighing 140 - 200 gm, each housed in separate compartments of a cage. All animals had free access to food and water. Different dosages of a compound were mixed with boiled fish and given to the animals at their habitual (7) feeding time. The mortality within

48 hours were recorded along with the toxic symptoms. Acute oral toxic dosage was found out by varying the amount of compound proportionately. The negligible amount of compound wasted by the animal during dieting was roughly accounted for in determining the dosage.

#### 10. PHYTOTOXICITY TESTS:

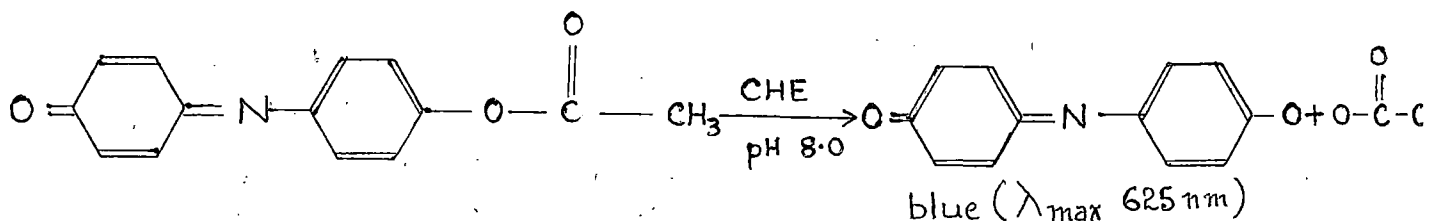
Phytotoxicity testing was conducted according to Eto et al. (8). Acetone solution of the compounds mixed with fixed amount of water containing 0.01% Triton - X was prepared. 5 ml of this aqueous suspension containing 500, 250 or 100 ppm of the compounds was poured into a petridish bottom covered with absorbent cotton. Ten seeds of wheat (Triticum sp UP 262 variety supplied by National Seed Corporation of India) were placed on the cotton and kept at room temperature (25 - 27°C) for five days. Occasionally 2 - 4 ml water was added in each petridish so that the seeds remained in moist condition. Each test was triplicated. Number of germination was counted after 5 days.

#### 11. ANTICHOLINESTERASE ACTIVITY:

The organophosphorus compounds have a common pharmacological property which is the ability to inhibit the activity of a group of enzymes, especially acetyl cholinesterase (AChE), involved in the hydrolysis of esters of choline. Since these enzymes present widely in insects and mammals, the organophosphorus compounds, used as insecticides also exhibit high mammalian toxicity.

## 11. (a) ANTIACHOLINESTERASE ACTIVITY IN HOUSEFLY - HEAD HOMOGENATE:

The acetyl cholinesterase inhibition of housefly head brie (HFACHE) has been measured by colorimetric method of Kramer and Gamson (9,10), using indophenyl acetate as an internal substrate-indicator in 0.05 M phosphate buffer of pH 8.0. The enzymatic reaction for indophenyl acetate is as follows:



The reaction mixture contained 5 ml buffer containing housefly head brie (1 ml of this solution contained 1 fly head) and 0.15 ml indophenyl acetate (total volume = 5.15 ml, conc. of indophenyl acetate  $\sim 10^{-5}$  M). The readings for control and sample were taken at 625 nm after exactly 30 minutes incubation.

### 11. (a) (1) REAGENTS:

1. Buffer solution : (0.05 M potassium dihydrogen phosphate):  
Clark and Lubs buffer of pH 8.0:

6.8 gm  $\text{KH}_2\text{PO}_4$  dissolved in 500 ml of water was mixed with 475 ml of 0.1 N NaOH solution and diluted to litre after the pH was adjusted to 8.0.

2. Indophenyl acetate + Working solution:

0.008 gm of indophenyl acetate when dissolved in 10 ml of

absolute methyl alcohol gave a  $3.3 \times 10^{-3}$  M solution so that the final concentration of the indophenyl acetate in the reaction vessel was always  $9.6 \times 10^{-5}$  M.

3. Glycerol solution:

10 ml of Glycerol was diluted to 100 ml with absolute alcohol.

4. Saline solution : 0.9%

9 gm of NaCl was dissolved in 1 litre distilled water

5. Salt solution:

2.03 gm of Manganous chloride and 2.15 gm of NaCl were dissolved in 250 ml of water.

6. Preparation of working solution of acetylcholinesterase from housefly-heads:

About 500 houseflies (Musca domestica) closed in a glass vessel was stored in a deep freeze for 1 - 2 hours. They were then transferred in a container with finely broken dry ice, removed individually from the container, decapitated with a shaving blade and forceps. 400 heads were combined with 2 ml of salt solution and 2 gms of washed sand in a prechilled with size No. 1 mortar. The heads were slowly ground, then transferred to 50 ml plastic centrifuge tube with one 3 ml aliquots of cold saline solution and two 5 ml aliquots of buffer solution. The head fragments were removed by centrifugation for

10 minutes at 10,000 r.p.m. in super speed centrifuge at 4°C. The supernatant liquid was decanted into a graduated cylinder and the fragmented heads were mixed with 10 ml of buffer solution and centrifuged again at 10,000 r.p.m. This extraction procedure was repeated twice. The supernatant solutions were combined and the volume was adjusted to 80 ml with the buffer solution so that each ml was equivalent to 5 fly-heads. This solution was stocked frozen in deep freeze. One ml of this solution was diluted to 5 ml with buffer solution so that each ml of the diluted solution contained single fly-head and used for each set of the experiment.

11. (a)(11) METHOD:

A series of 15 ml pyrex beakers (numbered 1,2,3..... etc.) containing different amounts of inhibitor (Viz, BD-10/ BD-11/etc.) in acetone along with one marked 'control' without inhibitors were arranged. 0.5 ml of glycerol solution was poured in each beaker including the 'control'. The acetone (in beaker 1,2,3 etc.) was removed by blowing cold air.

To the 'control' 5 ml of working enzyme - buffer solution was added and simultaneously the stop-watch was started. At the interval of exactly 2 minutes, 5 ml of the enzyme-buffer solution was added to each of the remaining beakers.

After exactly 30 minutes, 0.15 ml of the indophenyl acetate solution ( $3.3 \times 10^{-3}$  M) was added to the beaker marked 'control' and subsequently to each beaker of the series at the interval of 2 minutes and then kept to be incubated at 30°C. After incubation for

exactly 30 minutes the absorbances of 'control' and remaining solution were successively noted in the spectrophotometer (Carl - Zeiss Specol, ZV) at 625 nm with reference to enzyme-buffer (reagent blank) solution.

Calculation

$$\% \text{ inhibition} = \frac{\text{Absorbance (Control)} - \text{Absorbance (Sample)}}{\text{Absorbance (Control)}} \times 100$$

12. ANTICHOLINESTERASE ACTIVITY IN GOAT WHOLE BLOOD:

The method employed to determine the inhibition of the activity of acetyl cholinesterase in goat-whole blood by organophosphorus compounds was by colorimetric method of Kramer and Gamson<sup>(9,10)</sup>, using indophenyl acetate as an internal substrate indicator in 0.05 M phosphate buffer of pH 8.0. The reaction mixture contained 5 ml of enzyme buffer solution (4.8 ml phosphate buffer solution along with 0.2 ml goat whole blood) and 0.15 ml indophenyl acetate (total volume = 5.15 ml, concentration of indophenyl acetate in the reaction mixture  $\approx 10^{-5}$  M). The readings of 'control' and 'sample' were taken at 625 nm, after exactly 30 minute incubation.

12. (a) MATERIALS AND METHODS:

(1) Materials:

(a) Goat whole blood: 150 ml fresh blood was collected from goat and mixed with 15 mg ammonium oxalate (anticoagulating agent) in a 250 ml standard joint bottle and was shaken well. The bottle containing blood was then kept in the freezer at 0°C.

The other reagents e.g. (b) indophenylacetate (c) phosphate buffer solution and (d) glycerol solution were prepared just as the same as mentioned in 11 (a).

(11) Methods:

The anticholinesterase activity in goat whole blood was determined by the same method as describe in 11 (a), The only exception was that the contents of the beaker was filtered through a 4.25 cm whatman No-1 filter paper after exactly 29 minutes incubation and the absorbance of the filtrate of the enzyme buffer solution and % inhibition was calculated in the same way as describe in 11 (a).

13. CHEMICAL HYDROLYSIS:

The chemical hydrolysis studies were performed in 0.0095 M NaOH solution in 50 percent ethanol of pH 11.35 at 20°C. One ml of ethanol solution of the compounds was added with 9 ml of 0.0095 M NaOH solution (total volume 10 ml). The rate of hydrolysis was monitored by following the formation of nitro-saligenin anion at 410 nm in a Carl Zeiss Speckol ZV spectrophotometer. UV spectra of the alkylamidophosphorothionates and hydrolytic product were examined prior to Kinetic studies to show that overlap in relevant absorption peaks were not present. The concentration co-efficient value of 5 - nitro saligenin in the same 0.0095 M NaOH solution at the said wave lengths. The pseudo first order rate constants ( $K_{hyd}$ ) were determined by the least square regression analysis.

14. STUDIES ON FUNGICIDAL ACTIVITY USING SPORE GERMINATION METHOD:

14. (a) TEST ORGANISM:

1. Helminthosporium oryzae, Breda de Haan - causal organism of brown leaf spot disease of rice.
2. Verticillium albo-atrum, Reinke and Berthold - causal organism of stock rot disease of banana.
3. Aspergillus niger, Van Tiegh - causal organism of rot of stored grains.
4. Penicillium gensei, -causal organism of rot of stored grains.

14. (b) CULTURE MEDIUM:

Liquid medium : Malt solution.

40 gm. malt (Difco) extract was boiled in water till dissolved. The volume of the medium was then made upto 1 litre by adding water, pH of the medium was adjusted to 6.5 by adding NaOH solution. The medium was then sterilized at 15 p.s.i. for 20 minutes.

14. (c) SPORE GERMINATION METHOD:

Spore germination studies were made by using slide germination technique (11). Acetone solution of the compound was diluted

by sterile water in order to get the required amount of the compound in water suspension. Spore suspension of the fungus was prepared by flooding the surface of 12 to 15 days old culture with 5 ml sterile water; for A. niger and P. gansenii 0.01% agar solution was used in place of sterile water. Density of spore suspension in the final solution was determined by using a haemocytometer, and the final concentration was maintained at 20,000 spore/ml. Slides having spores in compound suspension were incubated at  $30^{\circ} \pm 1^{\circ} \text{C}$  and observation of the conidial germination were recorded after 24 and 48 hours. Three replication of each fungicidal concentration were maintained. A total of 300 spores from three replications were counted and percent inhibition of the spore germination was calculated against appropriate control.

15. FURTHER STUDIES ON FUNGICIDAL ACTIVITY ON OTHER FUNGI USING GROWTH INHIBITION METHOD:

Test organism:

1. Helminthosporium oryzae Breda de Haan;
2. Alternaria solani (BII and Mart) Jones and Grout - causal organism of early blight disease of potato.
3. Verticillium albo-atrum. Reinks and Berthhold;

The method used was stated previously [7 (c) - page - 109-110 ]

16. STUDIES ( in vivo ) OF FUNGICIDAL ACTIVITY OF SOME COMPOUNDS AGAINST  
H. oryzae ON RICE PLANT:

An isolate of H. oryzae was obtained from the Indian Agricultural Research Institute (New Delhi) and the culture was maintained on potato - dextrose-agar medium. Seeds of Dharial, a cultivar of rice susceptible to brown leaf spot disease, were used in the experiments. Seeds of this cultivar were grown at the Rice Research Station, Chinsurah (West Bengal). Naturally infected seeds of the same cultivar were also collected from the station and hand screened to give only spotted, infected seeds. Seventy to 75% of the spotted seeds yielded H. oryzae when grown in PDA.

16.(a) Determination of protectant activity on detached rice leaves:

Rice plants were grown in earthenware pots containing compost-enriched soil. When the plants were 40 days old, leaves of approximately the same age were washed, dried, removed from the plants and placed over moistened blotting paper taken in large petridishes. The required amount of compound in water suspension was sprayed on the leaves and allowed to dry for 15 to 20 minutes. Drops of conidial suspension of H. oryzae ( $8 \times 10^4$  spores/ml) prepared from 15 days old slant cultures were placed over the surface of the treated and untreated leaves. The petridishes were covered with polythene sheets and incubated at 21 - 23°C. After 48 hours production of lesion on leaves was recorded; percent inhibition of infection was calculated according to Chattopadhyaya and Bose (12)

to Chattopadhyaya and Bose .

16. (b) Protectant activity on rice seedlings:

Forty - day - old plants were sprayed almost to run off with water suspension of the compounds. Plants sprayed with water served as controls. After 12 hours, plants were inoculated by spraying with a conidial suspension of H. oryzae ( $8 \times 10^4$  conidia/ml). The inoculated plants were kept for 48 hours in a room in which high relative humidity and a temperature of 23 - 25° C were maintained. After incubation, the plants were placed in a greenhouse. The disease index was calculated after 6 days by using a modification of the method of Sinha and Trivedi<sup>(13)</sup>. The number and size of the spots appearing on leaves were recorded 6 days after incubation. The spots were graded into three size groups; small (upto 2.5 mm), medium (2.5 mm to 5 mm) and large (above 5 mm), with respective values of 0.25, 0.5 and 1.0 assigned to them. The number of spots in each size group was multiplied by the value assigned to it and the sum total of such values for all of the leaves gave the disease index for a plant<sup>(12)</sup>.

17. FURTHER STUDIES ON PHYTONKICIDITY OF SOME OF THE COMPOUNDS:

In this experiment above mentioned Dharial variety rice seeds have been used.

17. (a). Effect of treatment on rice seeds:

Well dried, cleaned seeds were soaked in acetone-water suspension of different compounds in two lots - one for two hours, and another for four hours respectively. The seeds were then dried in

open air. The seed germination were carried out on the next day in sterile petridishes containing moist filter paper. Hundred seeds were tested in each treatment and the germination percentage were observed and recorded. The germinated seeds were allowed to grow in petridishes for 7 days. The root and shoot length were recorded on 7th day. A parallel set of control was maintained always.

17. (b) Effect of treatment on rice plants:

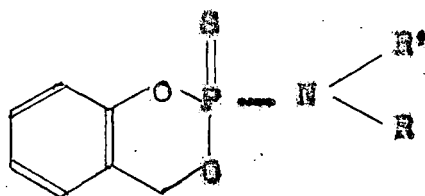
Thirty-day-old rice plants were sprayed almost to run off with water suspension of the compounds; the spraying was continued for consecutive 6 days (once daily). Each pot containing 10 plants were treated separately. Each experiment was done with five replications. A suitable control was maintained all along. The shoot length was measured before spraying (on 31 day) and after final completion of spraying (37th day).

B. RESULTS AND DISCUSSION

**B. RESULTS AND DISCUSSION:**

**1. SYNTHESIS**

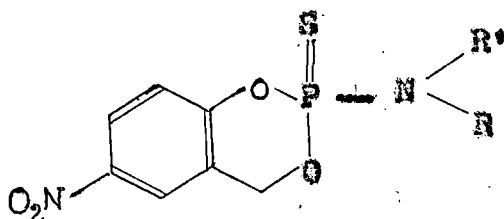
Only four saligenin cyclic phosphoramidothionates have so far been reported (14). The physical properties and percent yield of these compounds are given below.



Anido Group	Code No.	Yield percent	b.p. °C/mm Hg.
Methylamido	K - 35	20	120 - 123/0.2
Ethylamido	K - 37	--	Undistillable liquid
Dimethylamido	K - 36	27	118 - 122/0.2
Diethylamido	K - 38	13	110/0.2

All of them are liquids, and percent yield are low (13 - 27%)<sup>14</sup>.

We, however, succeeded<sup>to</sup> prepare the nitro-saligenin cyclic phosphoramidothionates (BD-10 to BD-18) in solid crystalline form with high yields (Table - 1).

Table - I

Code No.	Amido Group	Yield (%)	m.p. (°C)
BD - 10	Cyclohexylamido	70 - 90	125
BD - 11	Morpholine	60 - 80	149
BD - 12	Diethylamido	80 - 90	105
BD - 13	Dimethylamido	75 - 80	128
BD - 14	Isopropylamido	70 - 80	98
BD - 15	Pyrrolidino	80 - 90	134
BD - 16	Piperidino	80 - 90	130
BD - 17	Nonylamido	80 - 90	64
BD - 18	Heptadecylamido	80 - 90	70

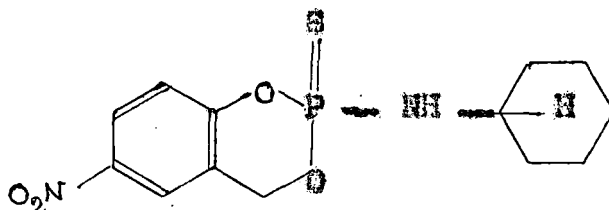
Attempts had also been made several times to prepare the 2-methylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide, and its ethylamido analog; unfortunately, in each case a viscous yellowish liquid was obtained. TLC of these liquids gave several spots indicating the presence of a number of compounds; the pure amidophosphorothionates could not be isolated from these liquid mixtures.

2. STRUCTURE DETERMINATION BY SPECTROSCOPIC METHOD:

The structure of the compounds have been determined by chemical analysis and IR, mass and NMR spectra. The analytical data have been presented in Section - A. The spectral data of only four compounds are given below:

2.(a) SPECTRAL DATA

(1) 2-Cyclohexylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide (BD-10):



IR (Fig. 1). (cm<sup>-1</sup>):

650 (m), 740 (m), 800 (s), 880 - 920 (s), 1020 (s), 1240 (vs),  
1340 (s), 1420 (m), 1480 (m), 1515 (s), 1584 (m), 1620 (w) and  
3300 (m).

Mass (Fig.2):

m/e	328 (M <sup>+</sup> ),	296,	295 (base peak),	279,	249
% RI	34.5,	43.5,	100	, 45.0,	33.0
m/e	230 ,	198	, 152		
%RI	35.0,	54.5	, 69.0		

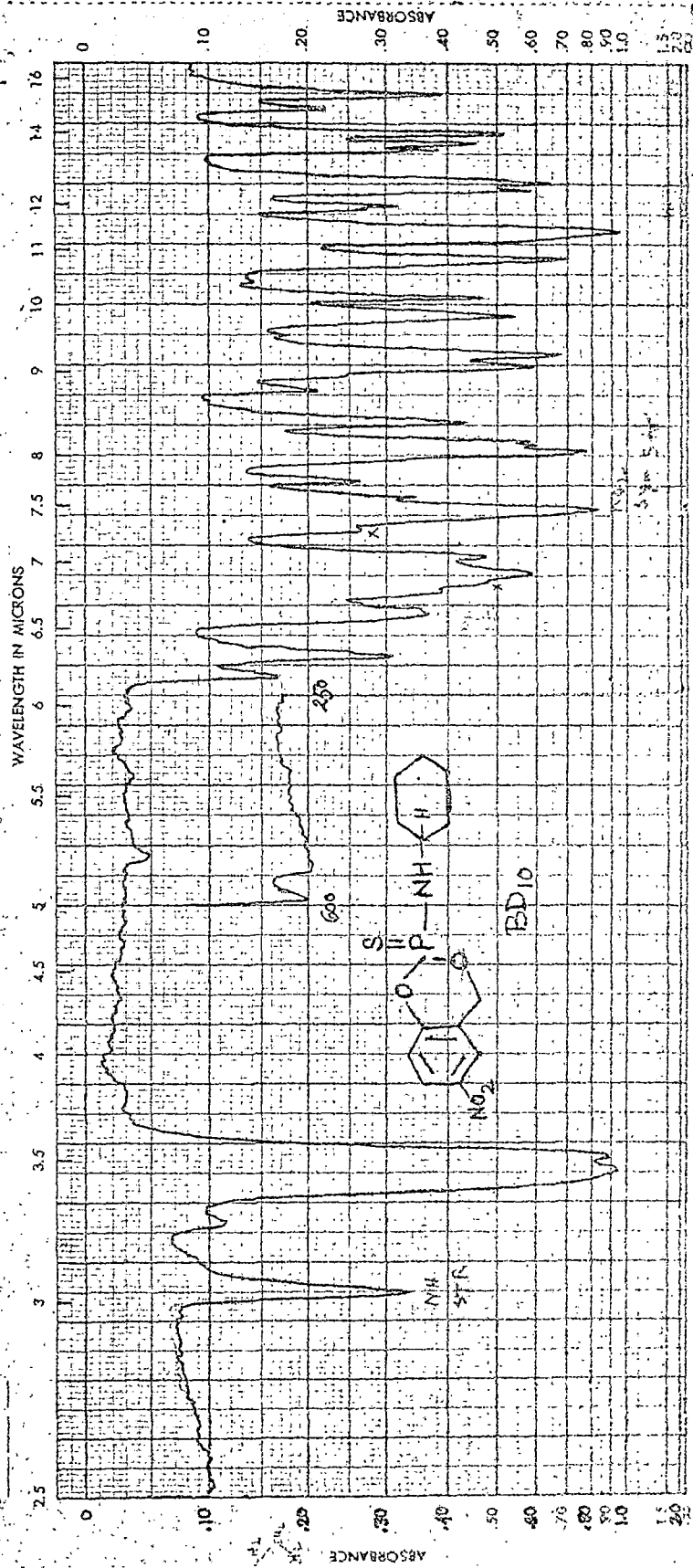


Fig. 1 IR spectra of 2-cyclohexylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide.

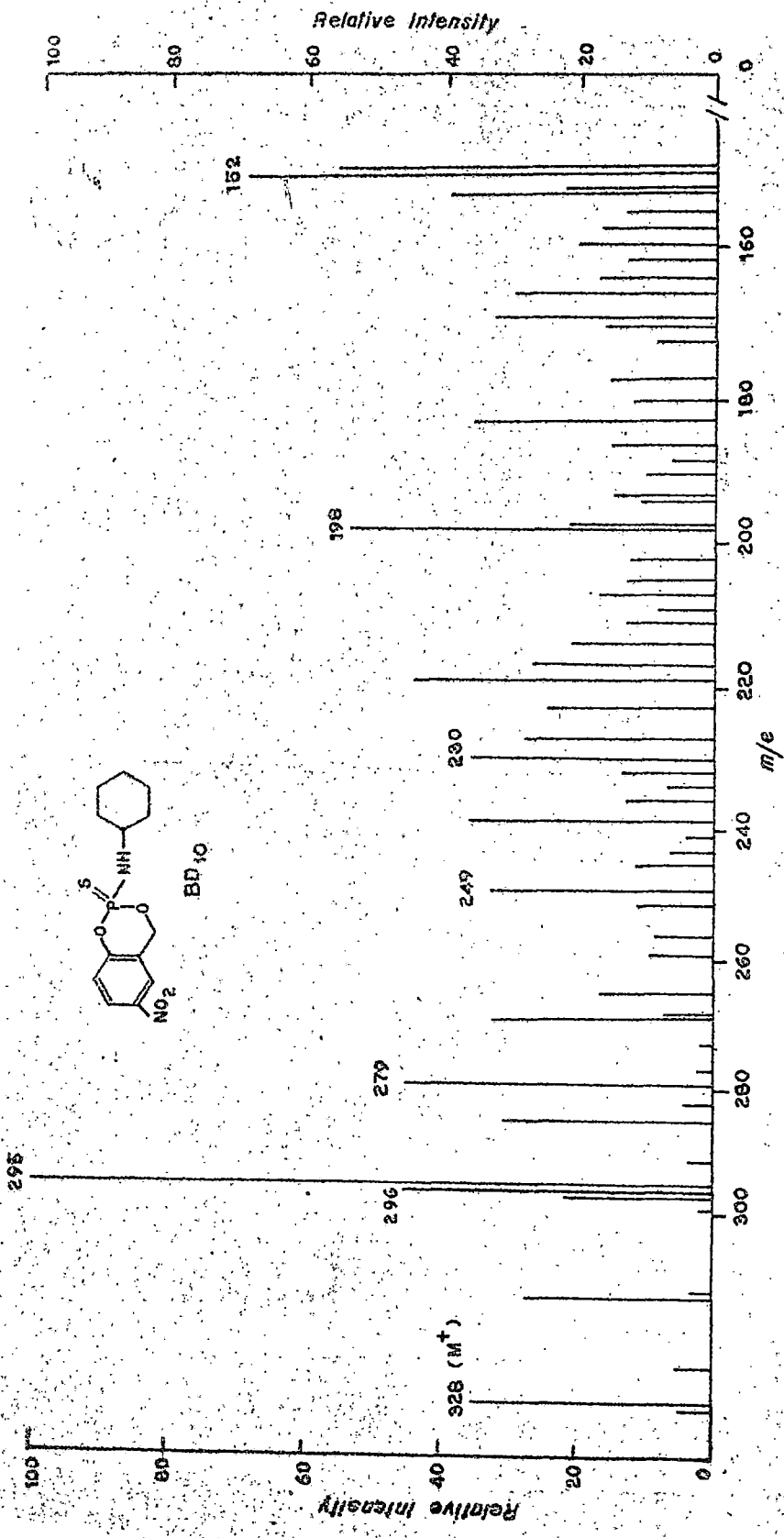
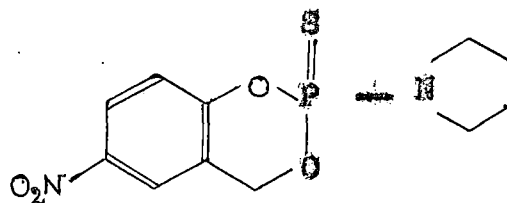


Fig. 2 Mass spectra of 2-cyclohexylamido-6-nitro-1,3,2-benzodioxaphosphorin-2-sulphide

$^1\text{H}$  NMR  $\delta$  ( $\text{CDCl}_3$ ) ppm (Fig.3):

- 1.25 (2H, multiplet,  $-\text{CH}_2$ -group at 4 position of the cyclohexylamin);
- 1.6 (4H, multiplet, two  $-\text{CH}_2$ -groups, at 3,3' position of the cyclohexylamin);
- 1.9 (4H, multiplet two  $-\text{CH}_2$ -groups at 2,2' position of the cyclohexylamin);
- 3.2 (1H, multiplet  $-\text{CH}<$  group);
- 3.6 (1H, multiplet  $-\text{P}-\text{NH}$  group);
- 5.45 (2H,  $\delta$ -line<sup>multiset</sup>  $-\text{CH}_2$ - group in dioxaphosphorin ring);
- 7.1 (1H, doublet,  $J = 8.5 \text{ Hz}$ , one aromatic hydrogen meta to nitro group);
- 8.05 (1H, doublet, one aromatic hydrogen ortho to both nitro group and  $-\text{CH}_2$ - group of dioxaphosphorin ring);
- 8.2 (1H, multiplet, remaining one aromatic hydrogen).

(ii) 2 - Pyrrolidino-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide  
(B9-15):



IR (Fig.4). ( $\text{cm}^{-1}$ ):

655 (s), 735 (m), 810 (s), 875 (s), 1030 (s), 1250 (s), 1340 (vs)  
and 1520 (vs).



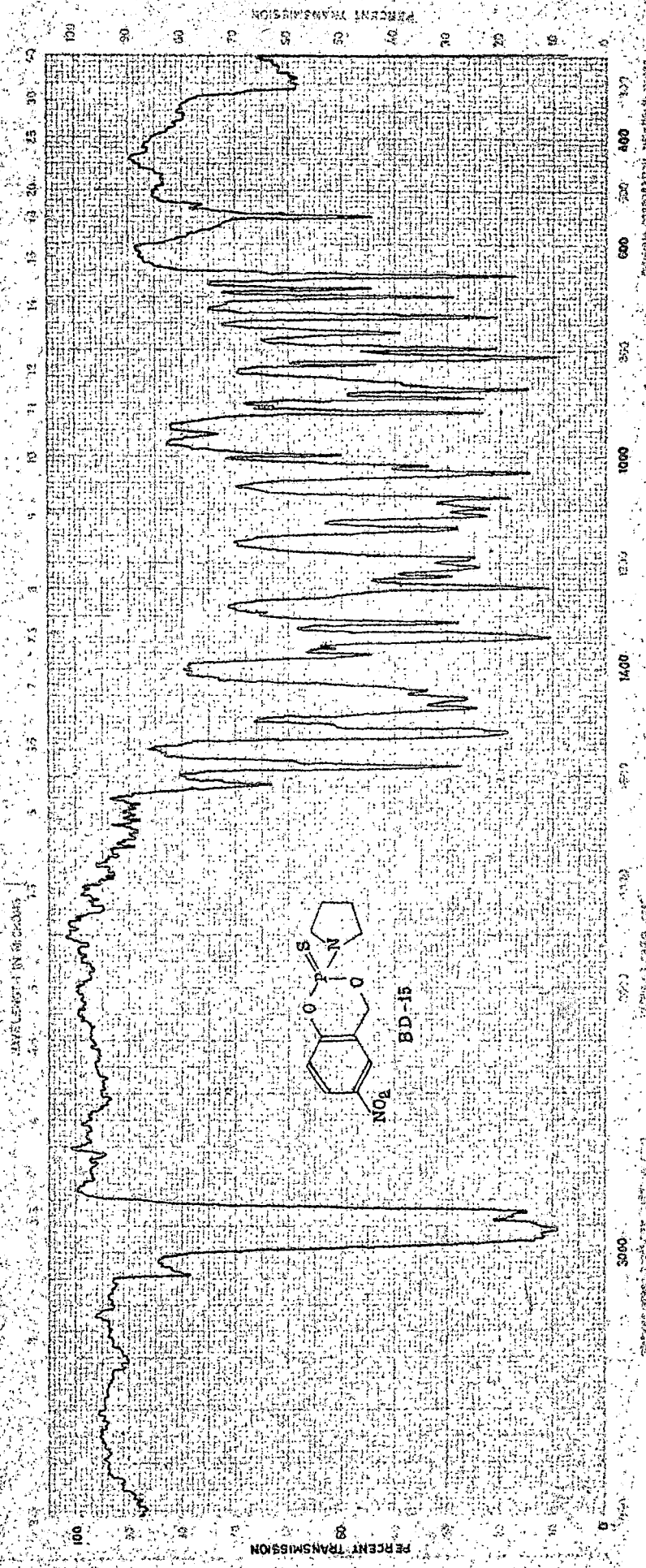


Fig. 4. IR Spectra of 2-Pyrrolidino-6-nitro-4M.1, 3, 2-benzodioxaphosphorin-2-sulphido.

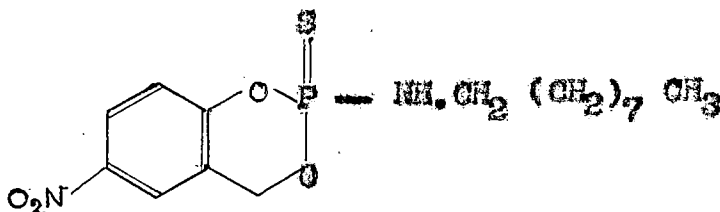
Mass (Fig. 5):

m/e	300 (M <sup>+</sup> )	, 267	, 198	, 148	, 116	, 70
%RI	21.4	, 100	, 25.7	, 11.4	, 80.0	, 35.7

<sup>1</sup>H NMR δ (CDCl<sub>3</sub>) ppm (Fig.-6):

- 1.62 - 1.95 (4H, multiplet, two -CH<sub>2</sub>-groups at 2, 2' positions of the pyrrolidine ring);
- 3.36 (4H, multiplet, two -CH<sub>2</sub>- groups adjacent to nitrogen);
- 5.26 and 5.65 (2H, -CH<sub>2</sub>- group in dioxaphosphorin ring);
- 7.05, 8.0 and 8.14 (due to aromatic hydrogens).

(111) 2-Nonylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide (BD.17):



IR. (Fig-7). (cm<sup>-1</sup>):

660 (s), 740 (m), 810 (s), 860 to 895 (s), 1030 (s), 1250 (s),  
1340 (s), 1520 (s), 1585 (m), 1620 (w) and 3310 (s)

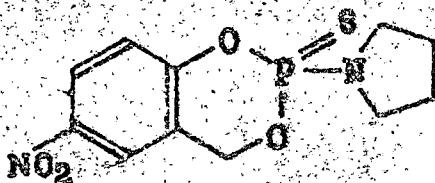
Mass (Fig-8)

m/e,	372 (M <sup>+</sup> )	, 339	, 259	, 230	, 198	, 152
%RI,	21.4	, 100	, 15.7	, 7.1	, 20	, 10

<sup>1</sup>H NMR δ (CDCl<sub>3</sub>) ppm (Fig-8A): 0.9, 1.2, 3.0, 3.4, 3.6, 5.5,  
7.05, 8.0, 8.2.

SPEKTRUM 23 VERDAMPFUNGSTEMPERATUR 120 GRAD  
 MOLEKUELPEAK: 300  
 MASSEN CHARAKTERISTISCHER IONEN:  
 267=300-SH

ANALYSE: 62907  
 STN HA 015 00  
 ST STEENKEN  
 MESSG:  
 AUSH : 25-MAR-81  
 AUSER: SCH



BD-15

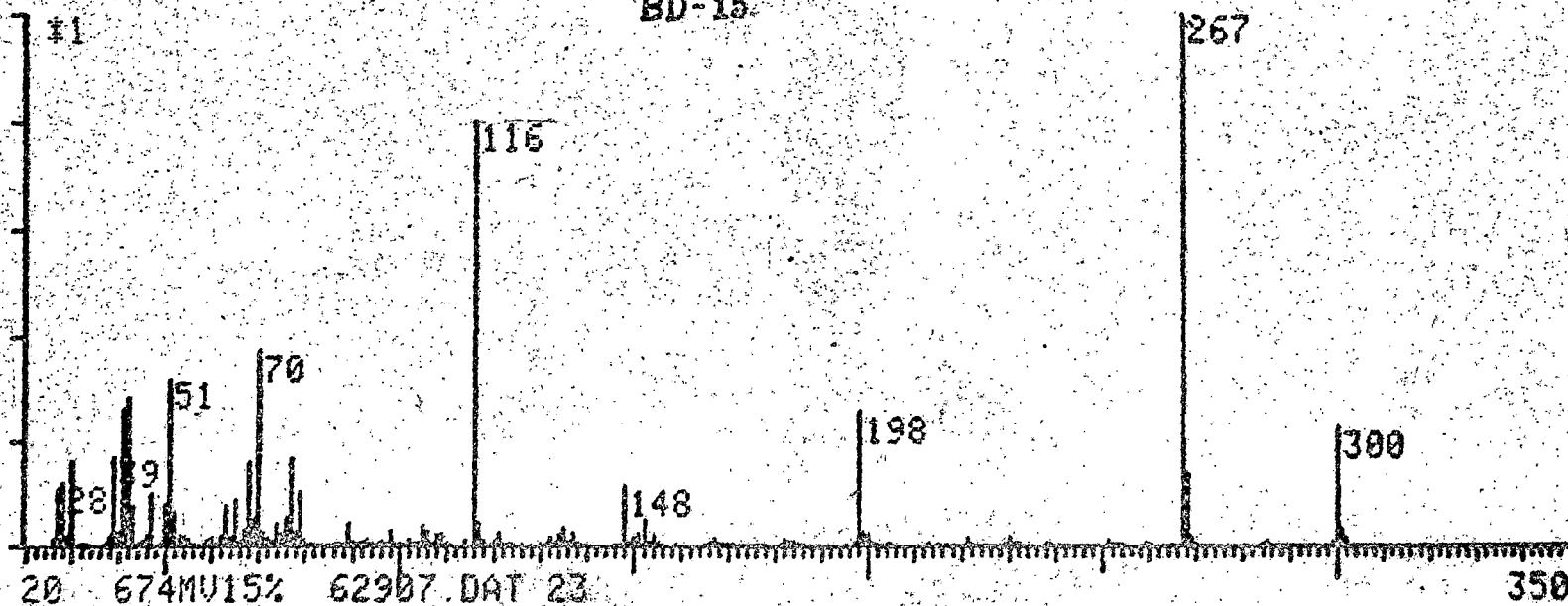
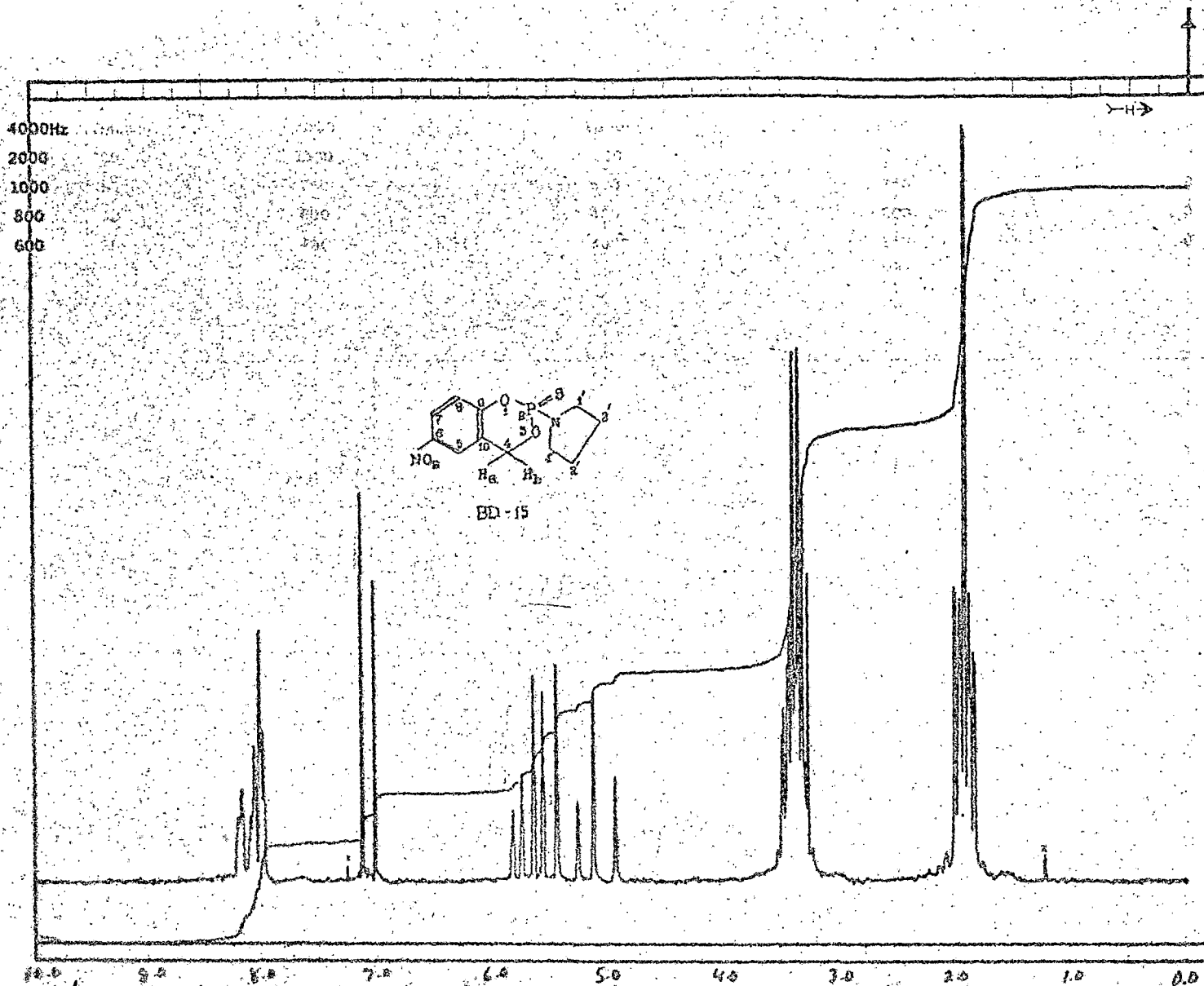


Fig. 5. Mass Spectra of 2-Pyrrolidino-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide.



FT-GOA SPECTRUM NO. 14992-H  
 OPERATOR gms DATE 1/18/81  
 NUCLEUS H FREQUENCY 79.542  
 SYNTHESIZER SETTING 77.2915  
 EXPERIMENT NAME \_\_\_\_\_  
 FILE NAME \_\_\_\_\_  
 SAMPLE BD-15

LOCK  INTERNAL  EXTERNAL  
 LOCK SIGNAL LO14  
 SPIN RATE 30 rps. TEMP 30 °C  
 INSERT 5 mm

ACQUISITION  
 SPECTRAL WIDTH (SW) 2000 Hz  
 NO. OF TRANSIENTS (NT) 50  
 ACQUISITION TIME (AT) 2.5 sec.  
 PULSE WIDTH (PW) 416 sec.  
 PULSE DELAY (PD) \_\_\_\_\_ sec.  
 DATA POINTS (DP) 114

TRANSMITTER OFFSET (TO) 77.2915  
 HIGH FIELD  LOW FIELD \_\_\_\_\_  
 RECEIVER GAIN (RG) 8

DECOUPLER MODE (DM) \_\_\_\_\_  
 DECOUPLER OFFSET (DO) \_\_\_\_\_  
 NOISE BANDWIDTH (NB) \_\_\_\_\_ kHz  
 ACQUISITION MODE (AM) \_\_\_\_\_

DISPLAY  
 SENS. ENHANCEMENT (SE) 1.5 sec.  
 WIDTH OF PLOT (WP) 795 Hz  
 END OF PLOT (EP) 0 Hz  
 WIDTH OF CHART (WC) 795 Hz  
 END OF CHART (EC) 0 Hz  
 VERTICAL SCALE (VS) 130  
 REFERENCE LINE (RL) (-606) 795

Fig. 6 <sup>1</sup>H NMR Spectra of 2-Pyrrolidino-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide.



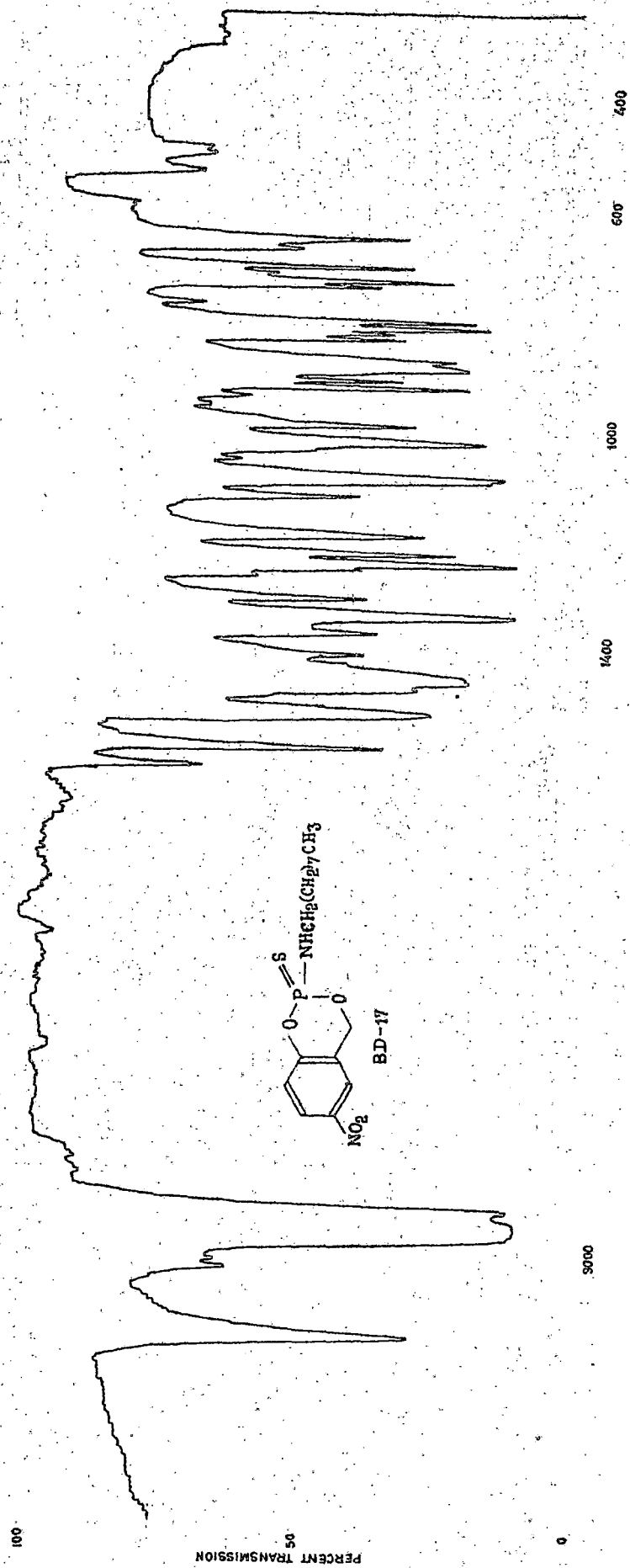


Fig. 7 IR Spectra of 2-Nonylamido-6-nitro-4H-1,3,2-benzoxaphosphorin-2-sulphide.

SPEKTRUM 45 VERDAMPFUNGSTEMPERATUR 140 GRAD  
 MOLEKUELPEAK: 372  
 MASSEN CHARAKTERISTISCHER IONEN:  
 339=372-SH

ANALYSE: 62895  
 STN NA 017 00  
 ST. STEENKEN  
 HESSG:  
 AUSH : 25-MAR-81  
 AUSMER: SCH

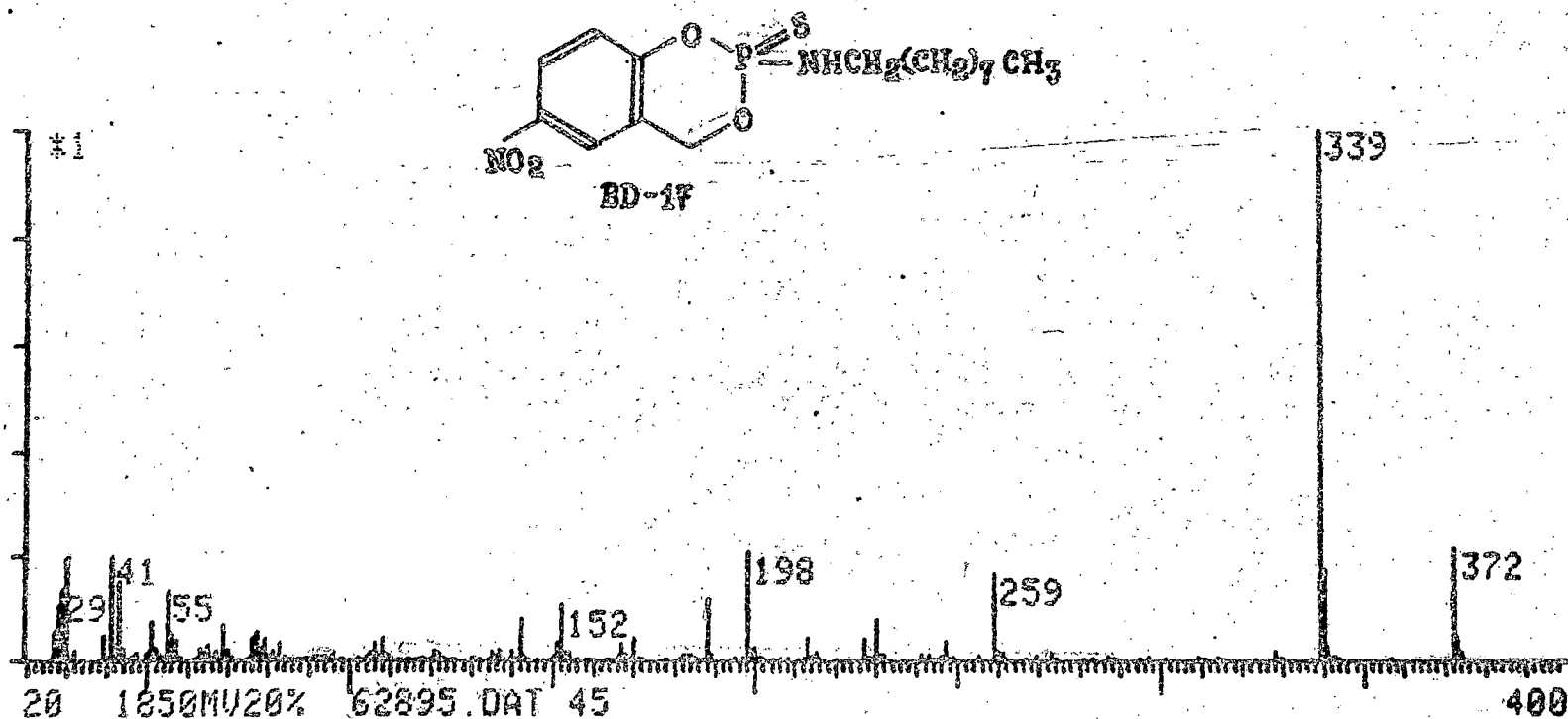


Fig. 8 Mass Spectra of 2-nonylsulphido-5-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide.

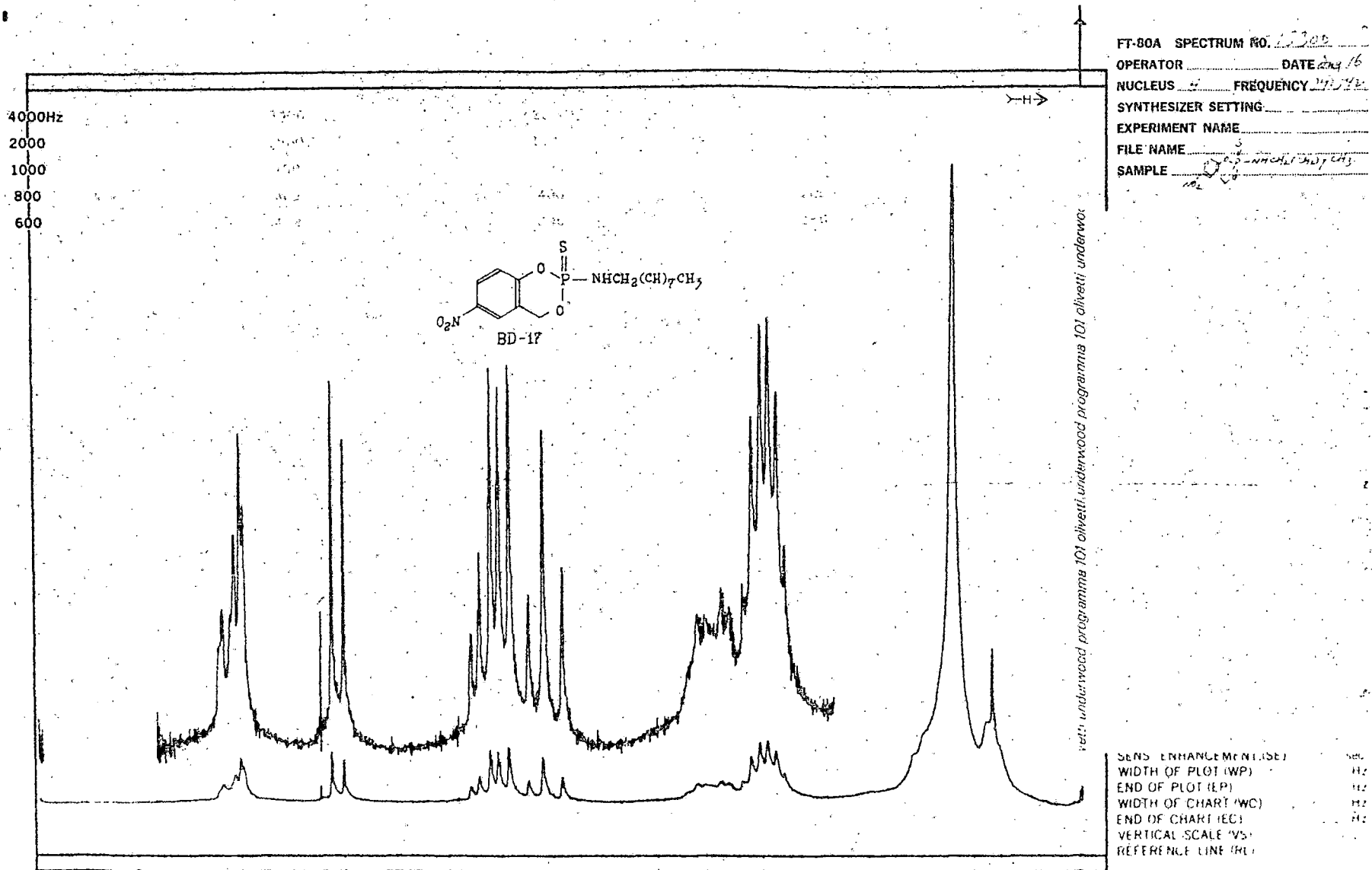
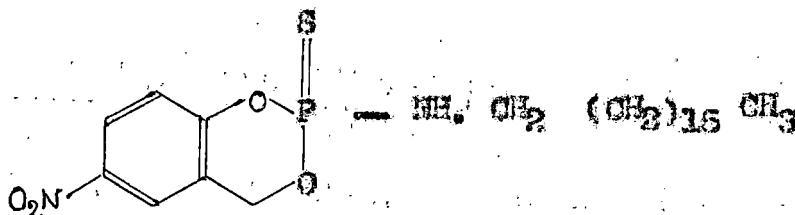


Fig. 8A. <sup>1</sup>H NMR Spectra of 2-Nonylamido-6-nitro-4H-1,3,2-benzodioxaphosphrin-2-sulphide.



(iv) 2 - Heptadecyl amido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide (BD-13):



IR (Fig. 9), (cm<sup>-1</sup>);

660 (m), 740 (m), 810 (s), 875 (s), 1030, 1245 (s),  
1345 (s), 1515 (s), 1590 (m), 1620 (w) and 3300 (s),

Mass (Fig - 10):

m/e, 484 (M<sup>+</sup>), 451, 300, 259, 230, 198, 152  
RI, 27.1, 100, 8.6, 29.3, 9.0, 28.6, 17.1

<sup>1</sup>H NMR δ (CDCl<sub>3</sub>) ppm (Fig-10A): 1.45, 3.15, 3.3 - 3.8, 5.5,  
7.05, 8.0, 8.15



SPEKTRUM 45 VERDAMPFUNGSTEMPERATUR 170 GRAD  
 MOLEKUELPEAK: 484  
 MASSEN CHARAKTERISTISCHER IONEN:  
 451=484-SH

ANALYSE: 62892  
 \*\*\*\*\*  
 STN NA 018 00  
 ST. STEENKEN  
 MESSG:  
 AUSH : 25-MAR-81  
 AUSHER: SCH

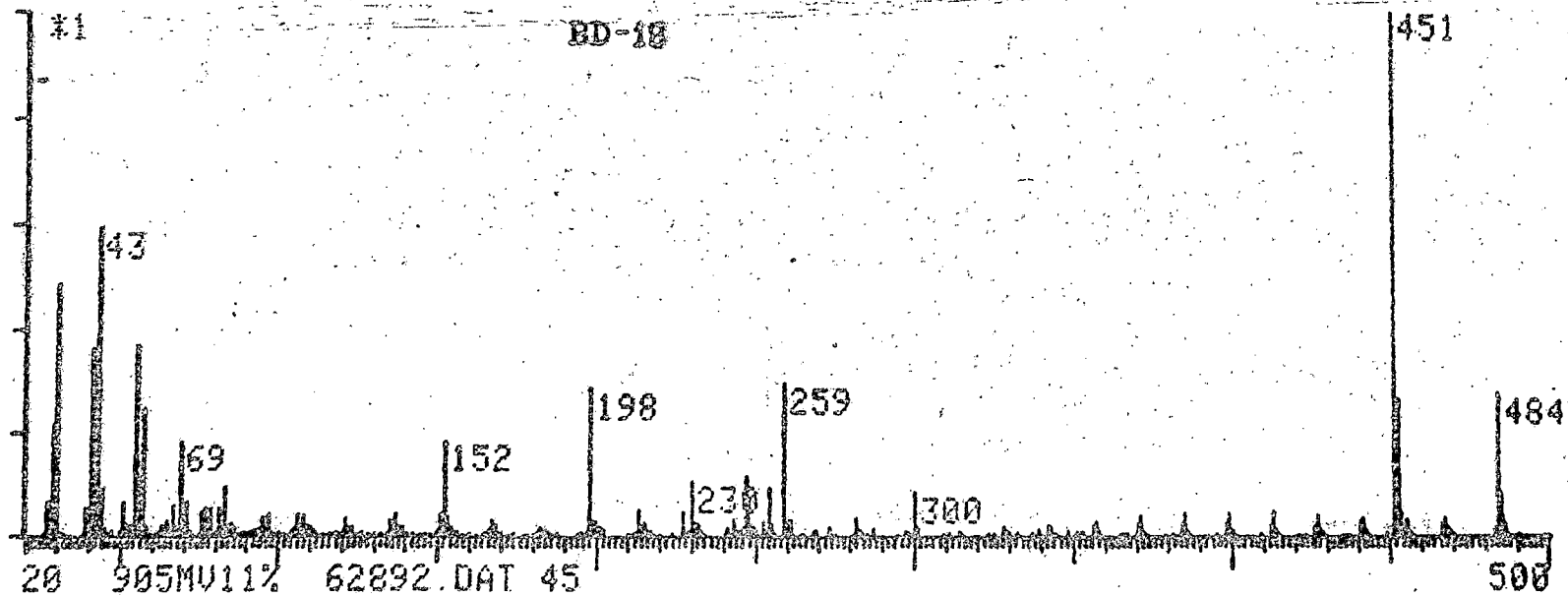
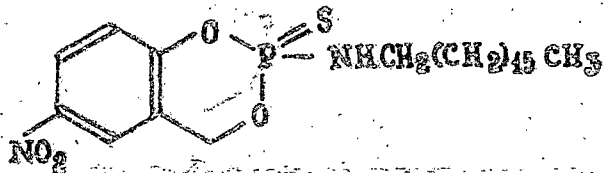


Fig.10 Mass Spectra of 2-Heptadecylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide.

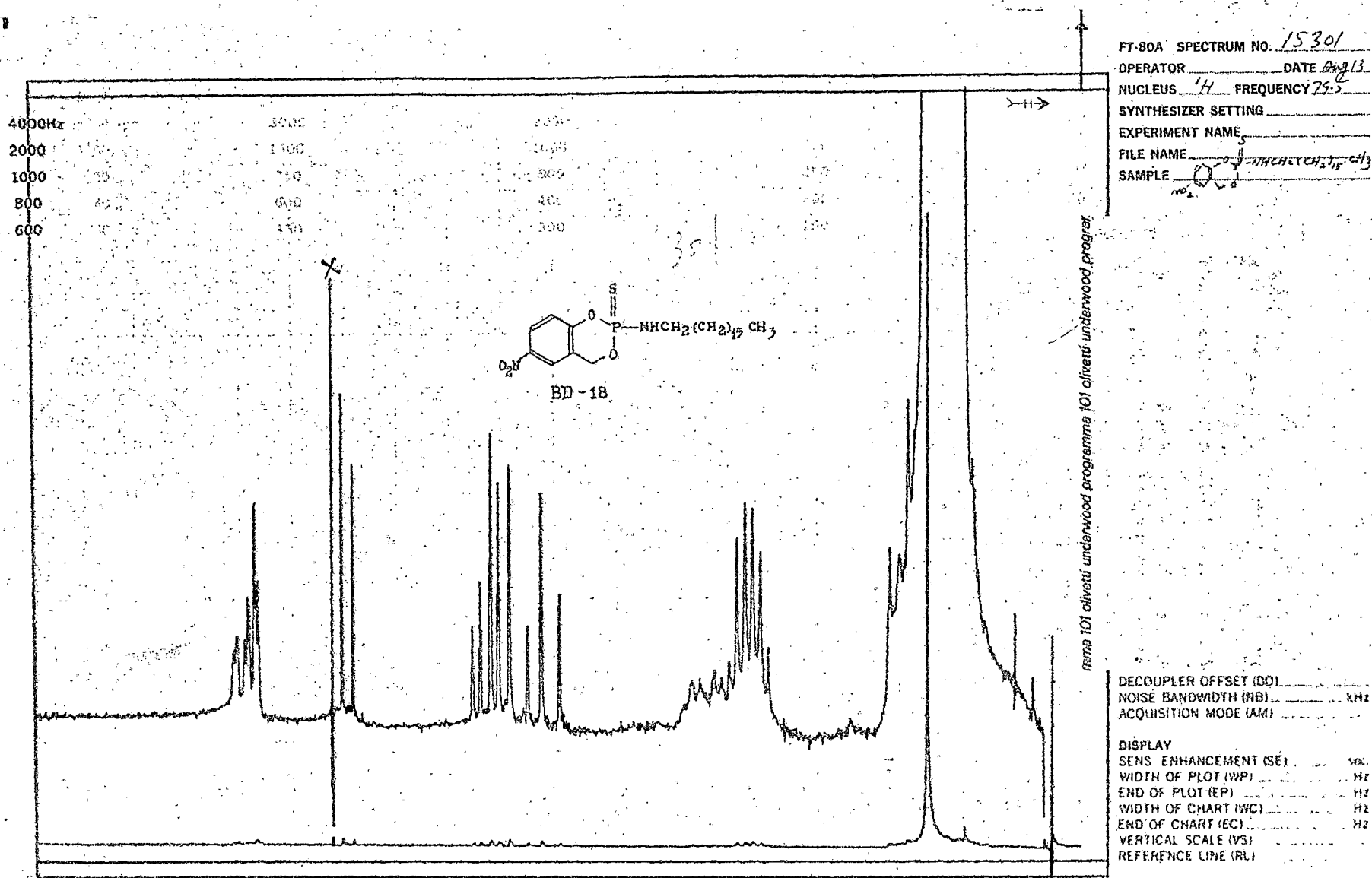


Fig. 10A. <sup>1</sup>H NMR Spectra of 2-Heptadecylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide.

2.(b) DISCUSSION ON SPECTRAL DATA:

The IR spectra of the compounds have been analysed according to Thomas <sup>(15)</sup>. The common IR bands for the compounds are summarized below:

1010 - 1030	$\text{cm}^{-1}$ (s), P - O - C (alkyl);
1235 - 1250	$\text{cm}^{-1}$ (vs), P - O - C (aryl);
880 - 920	$\text{cm}^{-1}$ (s), P - O - C (aryl);
1515 - 1520	$\text{cm}^{-1}$ (s), asym. str. of nitro group;
1340 - 1345	$\text{cm}^{-1}$ (s), sym. str. of nitro group;
800 - 820	$\text{cm}^{-1}$ , P = S (I);
640 - 660	$\text{cm}^{-1}$ , P = S (II);

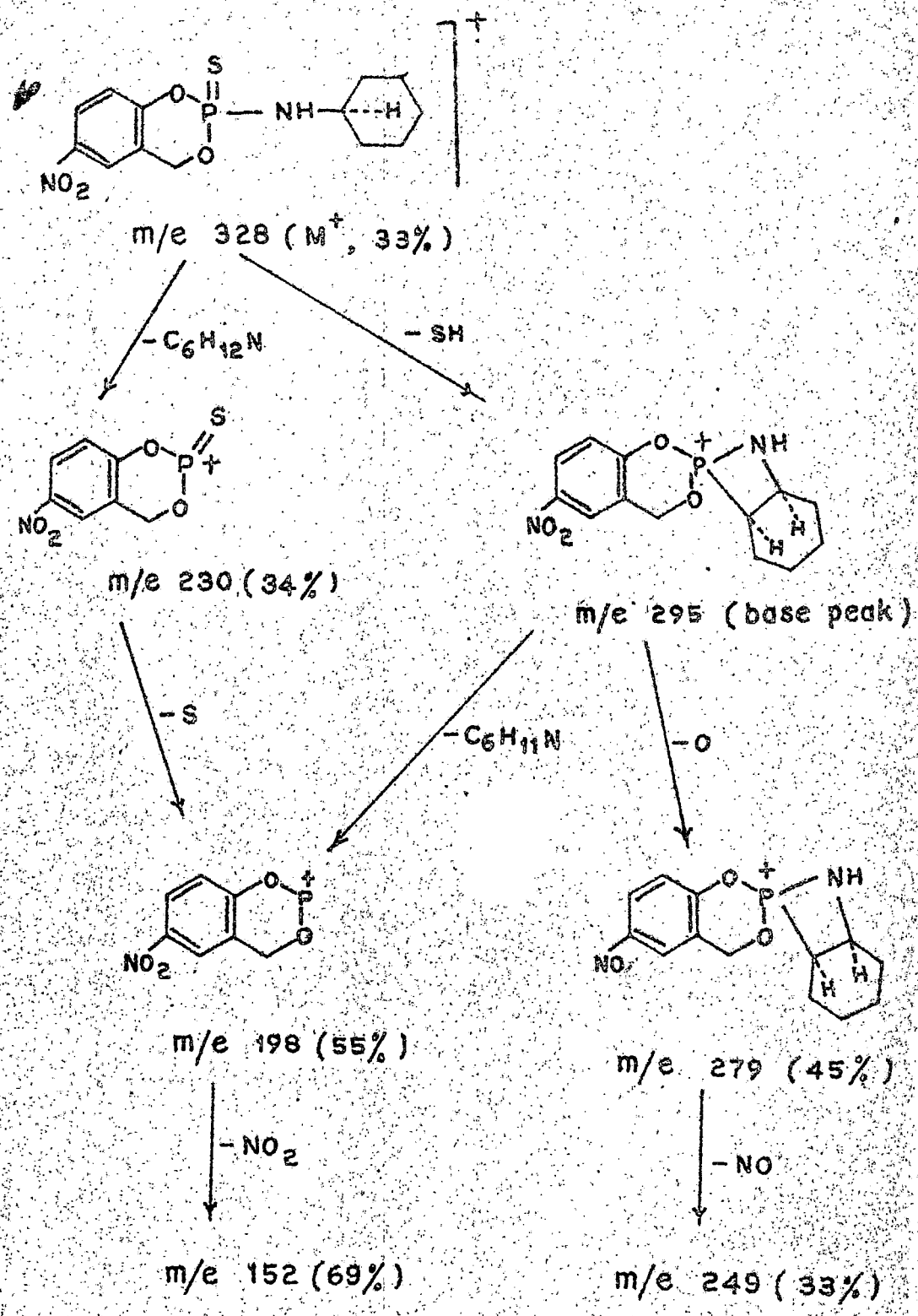
The thiono group is characterized by two IR absorption bands with frequencies in the normal ranges given by Thomas <sup>(15)</sup>, as both are not observed in phosphates; of these two, the lower frequency band is assigned to P = S (II) bond stretching vibration frequency. The origin of the higher frequency band P = S (I) is uncertain, but whatever its origin, its diagnostic value is beyond doubt. It is also observed that neither of two bands shows any systematic shifts which reflects changes in the inductive properties of the substituent; this is not unexpected if they do indeed arise from mixed modes. The P - O - C (alkyl) group is characterized by a strong absorption band whose frequency lies between 1010 - 1030  $\text{cm}^{-1}$ . While

the band due to P - O - C (aryl) group is found in the 1235 - 1250  $\text{cm}^{-1}$  region; this band is always accompanied by a second absorption band in the 880 - 920  $\text{cm}^{-1}$  region. Two bands present in the ranges 1515 - 1520  $\text{cm}^{-1}$  and 1340 - 1345  $\text{cm}^{-1}$  are due to asym. and sym. stretching of nitro group respectively. The bands present at 1620  $\text{cm}^{-1}$  and 1585  $\text{cm}^{-1}$  are due to "quadrant stretching" C = C vibration of the aromatic ring.

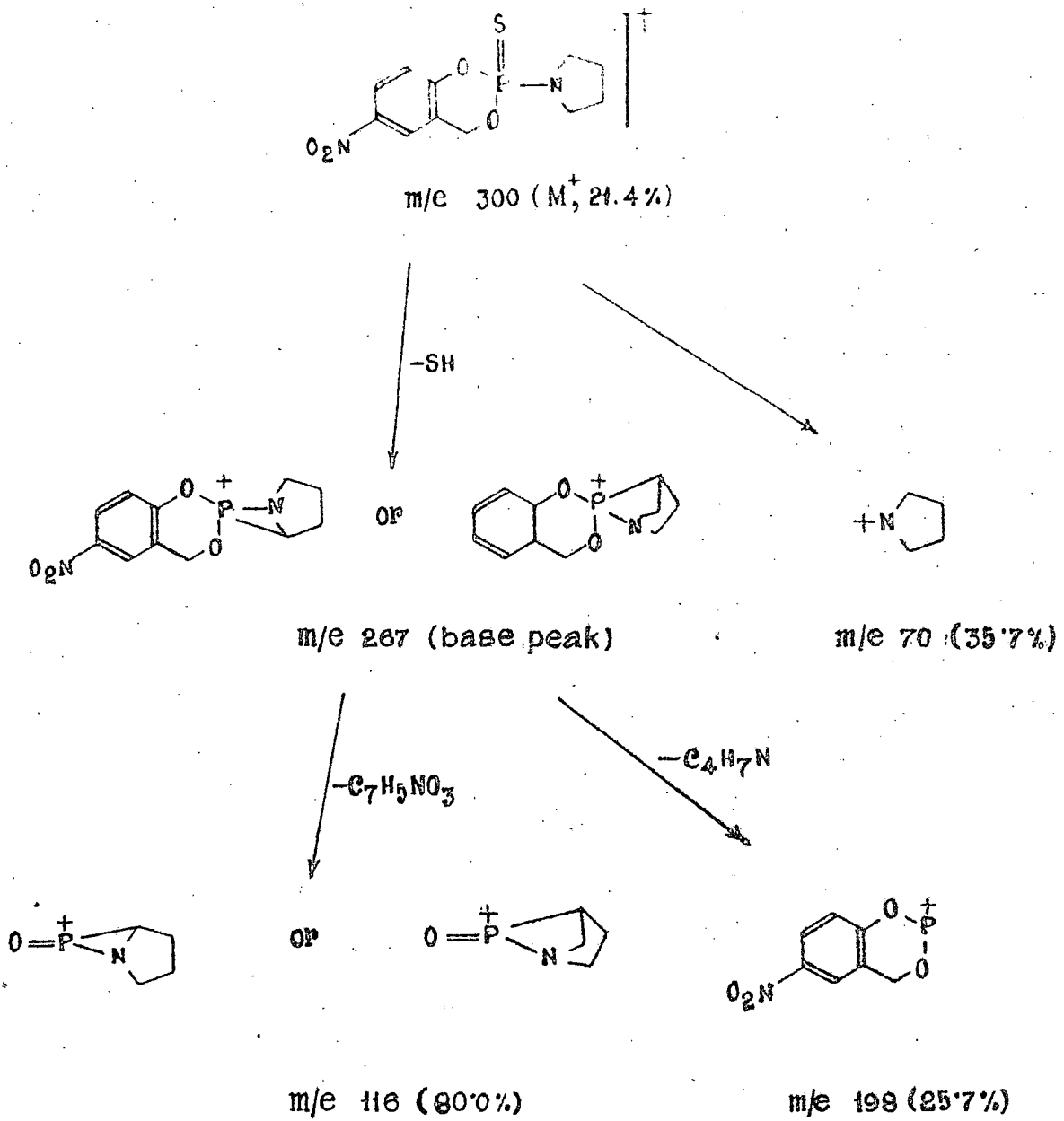
The mass spectra of these compounds have been analysed according to Cooks and Gerrard<sup>(16)</sup>. The alkylamidophosphorothionates show parent molecular ion peaks. Fragmentation by loss of 'SH' radical is important; all compounds (data for only four compounds have been presented) show an ion due to  $(M - SH)^+$ , and it is the base peak for all alkylamidophosphorothionates.

BD-10 shows m/e 295 ion as the base peak by the direct elimination of SH from the molecular ion peak (m/e 328, SRI 33, Scheme-I). The ion (m/e 230, SRI 34) is formed by the direct loss of  $\text{C}_6\text{H}_{12}\text{N}$  from the molecular ion. The ion (m/e 198, SRI 55) is formed by loss of 'S' from the ion m/e 230 and also by the elimination of  $\text{C}_6\text{H}_{11}\text{N}$  from the base peak ion. The ion (m/e 152, SRI 69) is formed by the elimination of  $\text{NO}_2$  from the ion m/e 198. The peak for m/e 279, SRI 45 is observed due to the elimination of 'O' from the base peak, and then the ion (m/e 249, SRI 33) is formed by loss of NO from the ion, m/e 279.

BD-15 shows m/e 267 ion as the base peak by the direct elimination of SH from the  $\text{M}^+$  ion peak (Scheme-II). The ion (m/e 116,

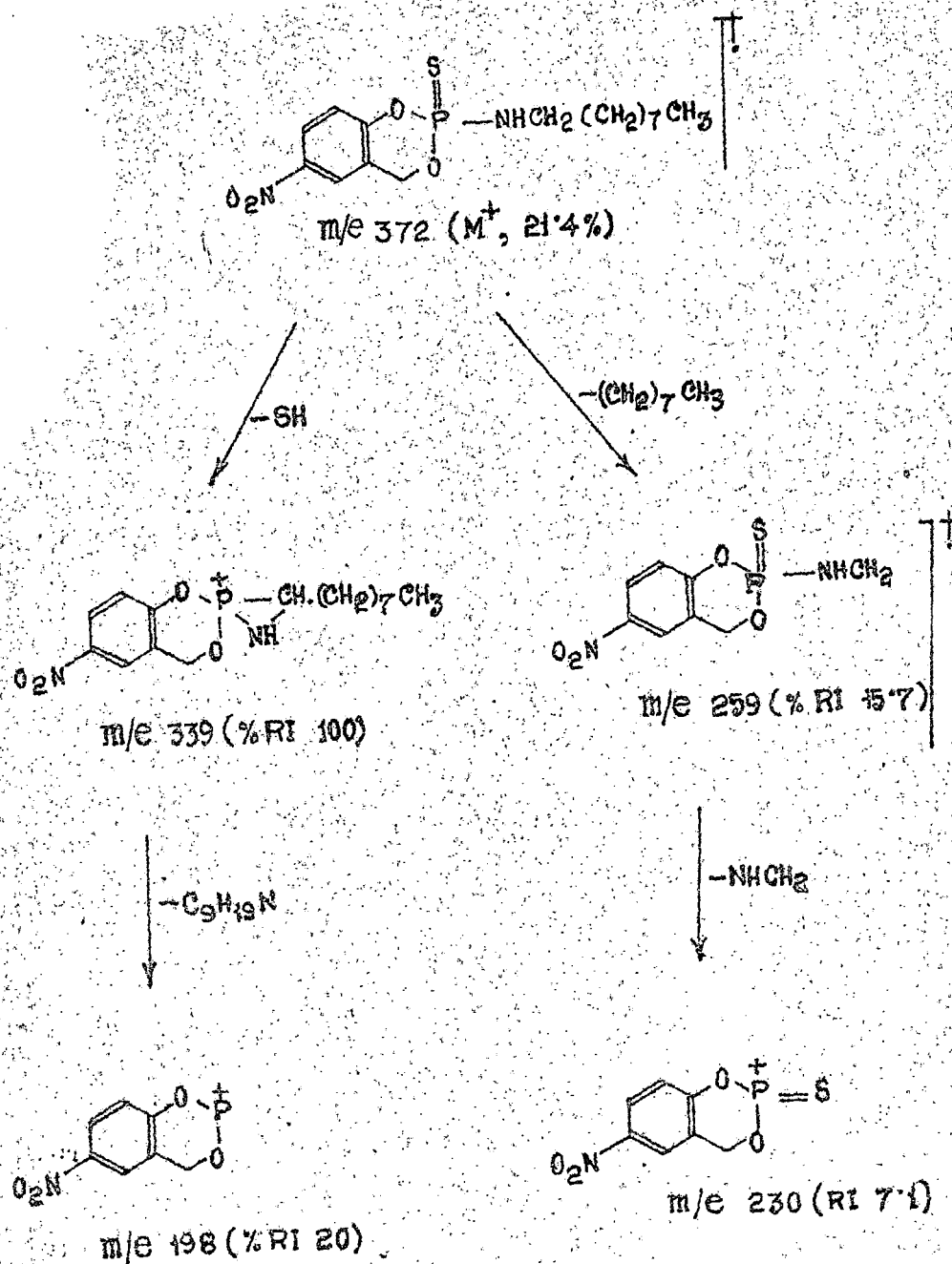


Scheme. I Mass fragmentation of BD<sub>10</sub>



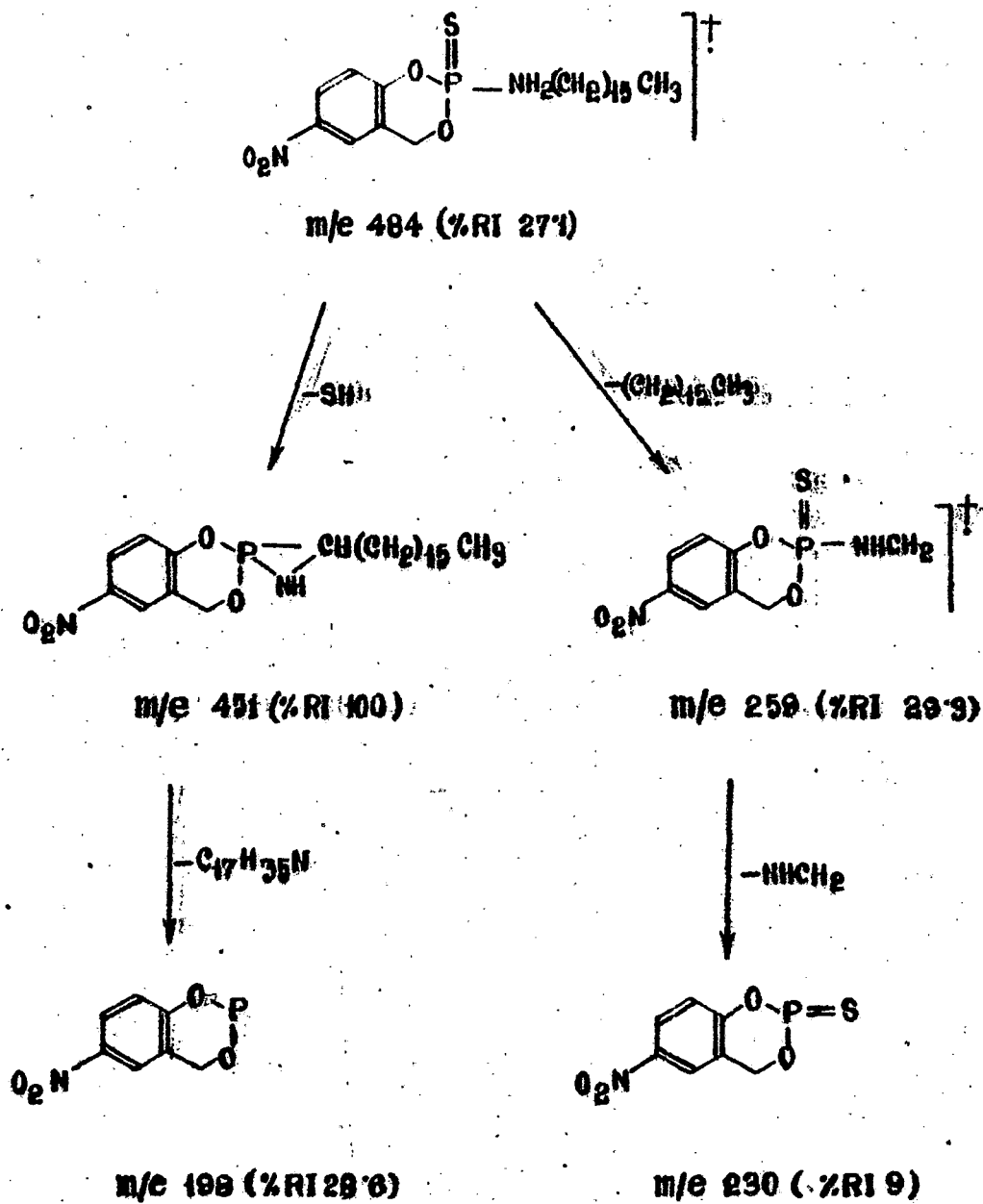
Scheme II

Mass fragmentation processes in BD-15



Scheme III

Mass fragmentation of BD-17



Scheme IV

Mass fragmentation processes in BD-18

§ RI 80) is formed by the direct loss of  $C_7H_5NO_3$  from the base peak ion; the ion ( $m/e$  198, § RI 25.7) is also formed by the direct loss of  $C_4H_7N$  from the base peak ion.

BD-17 shows  $m/e$  339 ion as the base peak by the direct elimination of  $SH$  from the parent molecular ion ( $m/e$  372, § RI 21.4, scheme III): The ion ( $m/e$  259, § RI 15.7) is formed by the direct loss of  $C_9H_{17}$  from the  $H^+$  ion peak; the ion ( $m/e$  198, § RI 20) is formed by the loss of  $C_9H_{19}N$  from the base peak ion. The ion ( $m/e$  230, § RI 7.1) is formed by the loss of  $CH_3N$  from the ion  $m/e$  259.

BD-18 shows  $m/e$  451 ion as the base peak by the direct elimination of  $SH$  from the molecular ion peak ( $m/e$  484, § RI 27.1) (scheme -IV). The ion ( $m/e$  198, § RI 23.3) is formed by the loss of  $C_{17}H_{35}N$  from the base peak ion. The ion ( $m/e$  259, § RI 29.3) is formed by the elimination of  $-(CH_2)_{15} \cdot CH_3$  from the molecular ion, and the ion ( $m/e$  230, § RI 9) is formed by the elimination of  $\cdot NH \cdot CH_2$  from the ion  $m/e$  259 (scheme-IV).

Finally the structure of different compounds has been settled by taking  $^1H$  NMR spectra, and analysis of spectral data has been given above [in 2(a)].

### 3. DISCUSSION ON FUNGICIDAL ACTIVITY (*P. oryzae* - Growth Inhibition):

The data (% inhibition) for in vitro growth inhibition studies against the blast disease of rice caused by *P. oryzae* are listed in Table - IA to IJ (Page-162-171). The  $ED_{50}$  values ( $\mu g/ml$ ) calculated

by least - squares regression analysis are given in Table - 1.

The results reveal that all compounds show good inhibitory effect on the growth of *P. oryzae*. At 72 hours the  $ED_{50}$  value increases in the following order:

BD-16 < BD-11 < BD-10 < BD-12 < BD-17 < BD-15 < BD-14 < BD-18 < BD-13

i.e. the dimethylamido compound (BD-13) is most effective, and its inhibitory effect can be comparable with that of Hinosan. At 48 hours both the dimethylamido and the isopropylamido (BD-14) compounds are most active; other compounds are also effective, but their activities are less than that of Hinosan. The  $ED_{95}$  values ( $\mu\text{g/ml}$ ) calculated from the least-square straight lines are presented in Table-2, and it can be observed that BD-13 and BD-14 are 2-times less active compared to Hinosan.

Table - 1

Fungicidal activity of different compounds against *Pyricularia oryzae*

Code name of the Compound	$ED_{50}$ in $\mu\text{g/ml}$			
	48 hours	72 hours	96 hours	120 hours
BD - 10	12.30	30.19	47.86	66.06
BD - 11	16.59	30.90	38.90	61.65
BD - 12	10.47	19.49	25.11	35.48
BD - 13	<5.00	5.88	10.96	21.87
BD - 14	<5.00	>12.50 <25.00	>12.50 <25.00	>25.00 >50.00
BD - 15	10.71	14.12	17.37	31.62
BD - 16	23.44	38.01	45.70	61.65
BD - 17	11.22	13.10	25.11	40.73
BD - 18	9.12	11.22	27.54	39.01
Hinosan (Standard)	<5.00	<5.00	<5.00	<5.00

Table - 2

Fungicidal activity of different compounds against Pyricularia oryzae.

Code name of the Compound	ED <sub>95</sub> in $\mu$ g/ml			
	48 hours	72 hours	96 hours	120 hours
BD - 10	60.25	275.42	537.03	616.59
BD - 11	132.03	229.03	251.13	339.04
BD - 12	69.18	134.89	151.35	194.93
BD - 13	>12.50 <25.00	25.70	43.97	134.89
BD - 14	>5.00 <12.50	>25.00 <50.00	>25.00 <50.00	>50.00 <100.00
BD - 15	58.83	91.20	93.32	169.82
BD - 16	154.83	275.42	281.83	363.07
BD - 17	63.09	123.02	158.48	194.93
BD - 18	45.70	132.03	147.91	177.82
Hinosan (Standard)	>5.00 <12.50	>12.50 <25.00	>25.00 <50.00	>25.00 <50.00

[Ref. Tables - 1A to 1J, page -162 to 171]

4. DISCUSSION ON INSECTICIDAL ACTIVITY;

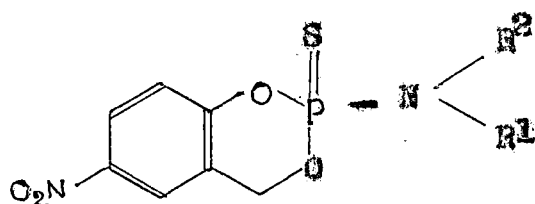
The insecticidal activity data of the compounds against Grasshoppers, Oxya nitidula are listed in Table - 3, and the results have been compared with that of salithion and the 2-methoxy - 6 - nitro-4H - 1,3,2 - benzodioxaphosphorin-2- sulphide (BD-8).

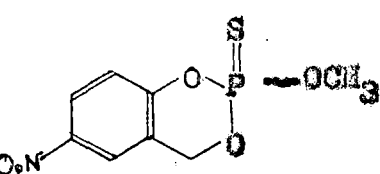
The data presented in Table - 3 reveal that all alkylamidophosphorothionates have less insecticidal activity than the methoxy compound (BD-8) and Salithion. The dimethylamido (BD-13) and the morpholino (BD-11) compounds have some insecticidal activity, but the other compounds are non-insecticidal. In whole series of nitro-saligenin cyclic alkyl/phenyl/alkylamidophosphorothionates synthesized in our Laboratory, only the methoxy compound (BD-8) shows insecticidal activity comparable with that of Salithion.

(17)

Table - 3

Insecticidal Activity on Grasshoppers (*Oxya nitidula*)



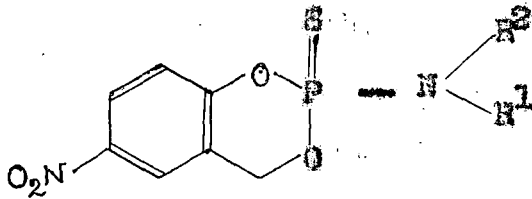
Amido Group	Code No.	Minimum Conc. (range ) for 100 percent morta- lity (LC <sub>100</sub> , μg/gm range).	Average LC <sub>100</sub> value (μg/gm)
Cyclohexylamido	BD - 10	>10	>10 (0%)
Morpholino	BD - 11	6 - 7	6.5
Dimethylamido	BD - 13	3 - 4	3.5
Isopropylamido	BD - 14	>10	>10 (0%)
Pyrrolidino	BD - 15	>10	>10 (0%)
Piperidino	BD - 16	>10	>10 (0%)
Nonylamido	BD - 17	>10	>10 (0%)
Heptadecylamido	BD - 18	>10	>10 (0%)
	BD - 8	< 0.5	< 0.5
Salithion	---	< 0.5	< 0.5

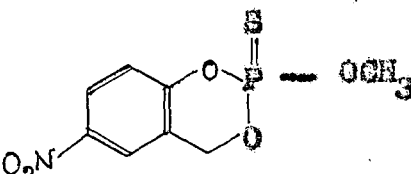
5. DISCUSSION ON ACUTE ORAL TOXICITY ON RAE:

The acute oral toxicity data ( $LD_{50}$ ) of the compounds (BD-10 to BD-15) on male rats are presented in Table-4, and the results have been compared with salithion and with the methoxy compound (BD-8). All compounds (BD-10 to BD-15) are less toxic than salithion. BD-12, BD-13, BD-15 and BD-8 have greater toxicity to rats compared to other nitro saligenin cyclic phosphoramidothionates. Before death the rats were found to suffer from acute respiratory trouble. In some cases, a fluid with blood-stain oozed out of nostrils and eyes of the animals. The decrease of spontaneous motor activity was occurred after 2 & 4 hours for all compounds except BD-15; in BD-15 it appeared after 60 - 72 hours.

Table - 4

LD<sub>50</sub> value of different compounds in Rats.



Amido Group	Code No.	LD <sub>50</sub> ( mg/kg)
Cyclohexylamido	BD - 10	> 250
Morpholino	BD - 11	> 250
Diethylamido	BD - 12	150 - 250
Dimethylamido	BD - 13	125 - 250
Isopropylamido	BD - 14	> 250
Pyrrolidino	BD - 15	150 - 250
	BD - 8	220 - 135
Salithion	--	102

6. DISCUSSION ON PHYTOTOXICITY (against wheat seed germination):

The effects of the alkylamidophosphorothionates on the germination of wheat seeds (*Triticum Spp*, U.P. 262 variety) are summarized in Table - 5. The results indicate that none of the compounds are phytotoxic upto 500 ppm concentration; however, BD-10, BD-11 and BD-17 show 90 percent germination at 500 ppm.

Table - 5

The effects of Alkylamidophosphorothionates on germination of wheat seed  
(Triticum Spp)

Code No.	Percent germination at different concentration		
	500 ppm	250 ppm	100 ppm
BD - 10	90	100	100
BD - 11	90	100	100
BD - 12	100	100	100
BD - 13	100	100	100
BD - 14	100	100	100
BD - 15	100	100	100
BD - 16	100	100	100
BD - 17	90	100	100
BD - 18	100	100	100

## 7. DISCUSSION ON ANTICHOLINESTERASE ACTIVITY:

The acetylcholinesterase inhibition data for housefly head homogenate (HFACH<sub>E</sub>) and goat whole blood (blood - ChE) are listed in Table 6A - 6J (page -172 to 181); the molar I<sub>50</sub> values calculated by least square programme are given below (Table - 6). The data for only five compounds have so far been taken.

Table - 6

Anticholinesterase activity on housefly head homogenate and goat whole blood.

Code No.	I <sub>50</sub> (M) × 10 <sup>5</sup> (HFACH <sub>E</sub> ) housefly.	I <sub>50</sub> (M) × 10 <sup>4</sup> (ChE) Goat whole blood
BD - 10	2.86	1.81
BD - 11	5.43	1.04
BD - 12	8.26	1.43
BD - 13	1.80	4.66
BD - 14	1.86	4.78

It has been observed that for any alkylamidophosphorothioates (BD-10 to BD-14), the housefly head acetylcholinesterase is more inhibited than the blood cholinesterase. For the housefly head acetylcholinesterase, the  $I_{50}$  value increases in the following order:



i.e., the antiacetylcholinesterase activity of the dimethylamido compound is highest, and that of the diethylamido-phosphorothionate is least. For the blood cholinesterase, the  $I_{50}$  value increases in the following order:



i.e. the anticholinesterase activity of the morpholino compound is most and that of the isopropylamidophosphorothionates is least.

#### 8. DISCUSSION ON CHEMICAL HYDROLYSIS:

The chemical hydrolysis of some of the compounds have been carried out in 0.0095 M NaOH (pH 11.85) at 410 nm; three sets of experiments in each case have been performed. The hydrolysis data for each compound at 410 nm along with the regression co-efficient value, and the value of  $K_{hyd}$  calculated thereof for a particular set are given in Table - 7A to 7E (page-182 to 186). The data have been summarized in the following table (Table - 7):

Table - 7

Hydrolysis of nitro saligenin cyclic alkylamidophosphorothionates  
(pH 11.85, temp. 20°C, solvent 50% ethanol).

Code No.	$K_{hyd}$ ( $\text{min}^{-1}$ )	Ref. Table No.	Average $K_{hyd}$ ( $\text{min}^{-1}$ )
BD - 11	$6.1413 \times 10^{-5}$	7A (page 182 )	$6.142 \times 10^{-5}$
BD - 13	$2.687 \times 10^{-5}$	7B (page 183 )	$2.88 \times 10^{-5}$
BD - 15	$1.7784 \times 10^{-5}$	7C (page 184 )	$1.763 \times 10^{-5}$
BD - 16	$1.2129 \times 10^{-5}$	7D (page 185 )	$1.162 \times 10^{-5}$
BD - 17	$20.266 \times 10^{-3}$	7E (page 186 )	$19.40 \times 10^{-3}$

The average hydrolysis constant recorded in Table - 7 show that the nature of the amide group in the exocyclic side chain influences the stability of the compounds to alkaline hydrolysis. The  $K_{hyd}$  increases in the following order:

$$BD - 16 < BD - 15 < BD - 13 < BD - 11 < BD - 17$$

i.e., the Piperidino compound (BD - 16) is most stable, and the nonylamido compound (BD - 17) is least stable to alkaline hydrolysis. The cyclic alkylphosphoramidothionates containing the disubstituted amide groups are extremely resistant to hydrolysis compared to the other compound having the monosubstituted amide group (nonylamido); probably the

the steric-interferences of the nonylamido group are less compared to that of the piperidino, pyrrolidino, morpholino and dimethylamido groups.

Contrary to alkaline hydrolysis in 0.0095 M NaOH these alkylamidophosphorothionates show a good deal of resistance to the hydrolysis at pH 7.7 in phosphate buffer. For example no detectable hydrolysis occurs even after thirtysix hours in case of dimethylamido, piperidino, pyrrolidino and morpholino compounds. However, the nonylamido compound shows slight hydrolysis at pH 7.7 in phosphate buffer; the rate of hydrolysis at pH 7.7 is less than that at pH 11.85. It may be concluded that the rate of alkaline hydrolysis is increased as the pH value increases from 7.7 to 11.8.

From the above studies it is clear that these alkylamido-phosphorothionates are almost non-insecticidal; however, they have good fungicidal activities, and have no phytotoxic properties. For these reasons further studies on fungicidal activities (both in vivo & in vitro) against some other pathogenic fungi, and also phytotoxic properties have been performed.

9. DISCUSSION OF FUNGICIDAL ACTIVITY ON OTHER FUNGI (Growth inhibition).

The data (5 inhibition) for in vitro studies against V. albo-atrum, A. solani and H. oryzae are listed in Tables - 8A to 8J. (page-187 to 196), 9A to 9I (page-197 to 205) and 10A to 10I (page-206 to 214) respectively. The  $ED_{50}$  and  $ED_{95}$  values ( $\mu$  g/ml) calculated by least square regression method are given in Tables - 8 to 10.

The results reveal that all compounds show inhibitory effect on the growth of V. albo-atrum, A. solani and H. oryzae. For V. albo-atrum at 48 hours the  $ED_{50}$  value increases in the following order:

BD-18 < BD-12 < BD-17 < BD-15 < BD-16 < BD-14 < BD-13 < BD-11 < BD-10

i.e., the cyclohexylamido compound (BD-10) is most effective, but its inhibitory effect is about 1.5 times less than that of Hinosan.

For A. solani at 72 hours the  $ED_{50}$  values increases in the following order:

BD-17 < BD-13 < BD-11 < BD-18 < BD-16 < BD-12 < BD-16 < BD-10.

i.e., the compound (BD-10) is most effective and its inhibitory effect can be comparable with that of Hinosan. The inhibitory effects of the compounds BD-12, BD-15 and BD-16 are nearly 1.5 times less than that of Hinosan.

For H. oryzae at 72 hours the  $ED_{50}$  values increases in the following order:

BD-11 < BD-12 < BD-13 < BD-17 < BD-10 < BD-15 < BD-16 < BD-14

i.e., the isopropylamido compound (BD-14) is most effective, and its inhibitory effect is nearly 1.6 times less than that of Hinosan.

To sum up, the dimethylamido compound is most effective against *F. oxysae*; the cyclohexylamido compound is most active against *V. albo-atrum* and *A. solani*; and, the isopropylamido compound is most effective against *H. oxysae*. But all of these compounds show less inhibitory effects than that of Hinosan.

Table - 8(1)

Fungicidal activity of different compounds against Verticillium albo-atrum

Code name of the Compound.	ED <sub>50</sub> in $\mu$ g/ml		
	24 hours	48 hours	62 hours
BD - 10	91.20	133.03	154.88
BD - 11	70.79	141.25	190.54
BD - 12	144.54	190.54	234.42
BD - 13	104.71	151.35	181.97
BD - 14	104.71	151.35	199.52
BD - 15	95.49	169.82	181.97
BD - 16	97.72	158.42	190.54
BD - 17	132.03	173.73	234.42
BD - 18	162.13	194.92	229.03
Hinosan (Standard)	>50.00	95.49	123.03

Table - 8 (2)

Fungicidal activity of different compounds against Verticillium  
alb-atrum.

Code name of the Compound	ED <sub>95</sub> in $\mu$ g/ml		
	24 hours	48 hours	62 hours
BD - 10	316.22	501.18	549.54
BD - 11	213.79	457.03	538.84
BD - 12	393.10	588.84	707.94
BD - 13	323.59	467.73	575.43
BD - 14	354.81	501.18	538.84
BD - 15	330.18	537.03	602.55
BD - 16	288.40	467.73	594.54
BD - 17	467.73	588.84	630.95
BD - 18	676.08	759.57	851.13
Hinosan (Standard) > 200.00		295.12	345.73

[ Ref. Tables 8A to 8J, page -187 to 196 ]

Table - 9 (1)Fungicidal activity of different compounds against Alternaria solani.

Code name of the Compound	ED <sub>50</sub> in $\mu$ g/ml			
	24 hours	48 hours	72 hours	96 hours
BD - 10	>50.00	75.85	120.22	147.91
BD - 11	<100.00	147.91	208.92	245.47
BD - 12	93.32	125.89	144.54	194.93
BD - 13	>50.00	134.89	229.03	281.83
BD - 14	<50.00	114.81	158.48	203.92
BD - 16	<50.00	89.12	128.82	169.92
BD - 17	100.00	173.78	257.03	288.40
BD - 18	50.00	77.62	136.20	239.88
Hinosan (Standard)	<50.00	60.25	104.71	169.82

[ Ref. Tables 9A to 9I, page -197 to 205 ]

Table - 9 (2)

Fungicidal activity of different compounds against *Alternaria solani*

Code name of the Compound	ED <sub>95</sub> in $\mu$ g/ml			
	24 hours	48 hours	72 hours	96 hours
BD - 10	>150.00	288.40	448.68	588.84
BD - 11	>150.00	575.43	616.59	676.03
BD - 12	399.02	363.07	436.51	549.54
BD - 13	>100.00	426.57	741.31	1000.00
BD - 15	>100.00	363.07	575.43	645.65
BD - 16	>100.00	323.59	426.57	575.43
BD - 17	>150.00	512.86	776.24	891.25
BD - 18	>150.00	245.47	588.84	776.24
Hinosan (Standard)	>100.00	177.82	309.02	448.68

[ Ref. Tables 9A to 9I, page -197 to 205 ]

Table - 10 (I)

Fungicidal effect of different compounds against Helminthosporium  
oryzae.

Code name of the Compound	ED <sub>50</sub> in $\mu$ g/ml			
	24 hours	48 hours	72 hours	96 hours
BD - 10	< 50.00	125.89	181.97	194.98
BD - 11	123.02	269.15	338.84	389.04
BD - 13	> 100.00	223.87	263.02	316.22
BD - 14	< 50.00	83.17	131.82	186.20
BD - 15	> 50.00	154.88	173.78	208.92
BD - 16	< 50.00	114.81	158.48	199.52
BD - 17	66.06	186.20	229.08	257.03
BD - 18	194.98	261.18	316.22	363.07
Hinosan (Stan- dard)	39.81	63.09	81.28	93.32

⌈ Ref. Tables 10A to 10I, page-206 to 214 ⌋

Table - 10 (2)

Fungicidal effect of different compounds against *Helminthosporium*  
*oryzae*.

Code name of the Compound.	ED <sub>95</sub> in $\mu$ g/ml			
	24 hours	48 hours	72 hours	96 hours
BD - 10	>100.00	500.00	575.43	602.55
BD - 11	562.34	977.23	1122.01	1548.81
BD - 13	>150.00	724.43	851.13	954.99
BD - 14	>100.00	263.02	407.38	630.95
BD - 15	>150.00	500.00	524.80	616.59
BD - 16	>100.00	549.54	602.55	752.57
BD - 17	346.73	524.80	602.55	707.94
BD - 18	524.80	707.94	891.25	1023.29
Hinosan (Standard)	186.20	213.77	269.15	309.02

[Ref. Tables 10A to 10I, page - 206 to 214]

10. DISCUSSION ON FUNGICIDAL ACTIVITY (Spore germination).

The data (% inhibition) for in vitro spore germination inhibition against A. niger & P. gansenii (Stora Fungi), and H. oryzae & V. albo-atrum are listed in Tables - 11A to 11I (page - 215 to 223 ), 12A to 12I (page - 224 to 232 ), 13A to 13I (page - 233 to 241 ) and 14A to 14I (page - 242 to 250). respectively. The  $ED_{50}$  and  $ED_{95}$  values are given in Tables 11 to 14.

The results reveal that all compounds are effective on spore germination inhibition of A. niger; P. gansenii; H. oryzae and V. albo-atrum.

For A. niger at 48 hours the  $ED_{50}$  value increases in the following order:

$BD-16 < \text{Hinosan} < BD-10 < BD-11 < BD-12 < BD-13 = BD-15 < BD-14 < BD-17$   
i.e., the inhibitory effects of all compounds except BD-16; are greater than that of Hinosan; the nonylamido compound (BD-17) is the most effective one.

For P. gansenii at 48 hours the  $ED_{50}$  value increases in the following order:

$BD-16 < BD-14 < BD-11 < BD-12 < BD-10 < \text{Hinosan} < BD-15 = BD-17 < BD-13$   
i.e., the dimethylamido compound (BD-13) is most effective, and its inhibitory effect is about 1.5 times greater than that of Hinosan. BD-15 and BD-17 are also active (activity is greater than Hinosan).

For H. oryzae at 48 hours the  $ED_{50}$  value increases in

the following order:

BD-16 < BD-15 < BD-13 < BD-11 < BD-18 < BD-10 < BD-17 < BD-14 < Kinosan  
i.e., the isopropylamido compound is most effective, but its inhibi-  
tory effect is about 1.5 times less than that of Kinosan.

For V. albo-atrum at 48 hours the ED<sub>50</sub> value increases  
in the following order:

BD-10 < BD-18 < BD-16 < BD-15 < BD-11 < BD-17 < BD-14 < Kinosan < BD-13

i.e., the dimethylamido compound is the most active, and its inhibitory  
effect is greater than that of Kinosan. Other compounds are less ac-  
tive.

To sum up, the nonylamido compound is most effective  
against A. niger; the dimethylamido compound is most active against  
P. versicolor and V. albo-atrum; and the isopropylamido compound is most  
effective against H. oryzae.

Table - 11

Fungicidal activity of some amide compounds against *A. niger* and *P. ganseri* (Spore germination).

Code name of the Compound.	ED <sub>50</sub> values in $\mu$ g/ml			
	<i>Aspergillus niger</i>		<i>Penicillium ganseri</i>	
	24 hours	48 hours	24 hours	48 hours
BD - 10	1096.47	1445.43	912.01	1288.24
BD - 11	1258.92	1412.53	1513.56	1819.70
BD - 13	776.24	1230.26	630.95	891.25
BD - 14	794.32	1174.89	1445.43	1862.08
BD - 15	870.96	1230.26	812.83	1023.29
BD - 16	1548.81	1905.46	1412.53	1949.84
BD - 17	831.76	1000.00	758.57	1023.29
BD - 18	1096.47	1318.25	1096.47	1412.53
Hinosan (Standard)	1067.12	1862.08	891.25	1202.26

[Ref. Tables 11A to 11 I, page - 215 to 223 ]  
12A to 12 I, page - 224 to 232.

Table - 12

Fungicidal activity of some amide compounds against *A. niger* and *P. gossypii* (Spore germination)

Code name of the compound	ED <sub>95</sub> values in $\mu$ g/ml			
	<i>Aspergillus niger</i>		<i>Penicillium gossypii</i>	
	24 hours	48 hours	24 hours	48 hours
BD - 10	2454.70	3530.78	1862.08	2511.88
BD - 11	1995.26	2630.26	2951.20	3235.93
BD - 13	2290.85	2818.38	1230.26	1549.81
BD - 14	2398.83	3019.95	2884.03	4168.69
BD - 15	1819.70	2754.22	1778.27	2344.22
BD - 16	3162.27	3981.07	3162.27	3801.89
BD - 17	1479.10	1905.46	1584.89	2137.96
BD - 18	2754.22	3388.44	2344.22	2630.26
Hinosan (Standard)	3811.31	5129.61	5187.76	2454.70

11 A to 11 I, page - 215 to 223  
 [Ref. Tables 12A to 12I, page - 224 to 232]

Table - 13

Fungicidal activity of some amido compounds against *H. oryzae* and *V. albo-atrum* (Spore germination).

Code name of the Compound	ED <sub>50</sub> values in $\mu$ g/ml			
	<i>Helminthosporium oryzae</i>		<i>Verticillium albo-atrum</i>	
	24 hours	48 hours	24 hours	48 hours
BD - 10	812.83	1023.29	1737.80	2691.53
BD - 11	1047.12	1258.92	1230.26	1445.43
BD - 13	1071.51	1286.24	630.95	977.23
BD - 14	776.24	831.76	776.24	1318.25
BD - 15	1071.51	1318.25	1230.26	1621.81
BD - 16	1122.01	1412.53	1348.96	1778.27
BD - 17	758.67	977.23	1623.29	1345.96
BD - 18	977.23	1071.51	1698.24	1995.26
Kinosem (Standard)	338.84	588.84	467.73	1122.01

[ Ref. Tables - 13A to 13I, page - 233 to 241 ]  
14A to 14I, page - 242 to 250

Table - 14

Fungicidal activity of some amido compounds against *H. oryzae* and *V. albo-atrum* (Spore germination).

Code name of the compound	ED <sub>95</sub> values in $\mu\text{g/ml}$			
	<u><i>Helminthosporium oryzae</i></u>		<u><i>Verticillium albo-atrum</i></u>	
	24 hours	48 hours	24 hours	48 hours
BD - 10	1513.56	1659.68	4570.88	6918.30
BD - 11	1698.24	1905.43	2290.86	2398.83
BD - 13	2137.96	2187.76	1253.92	1819.70
BD - 14	1230.26	1380.38	1584.89	3162.27
BD - 15	2398.83	2630.26	3019.95	3543.13
BD - 16	2454.70	2570.39	3235.93	3388.44
BD - 17	1258.92	2041.73	2290.86	2570.39
BD - 18	1698.24	1778.27	3715.35	4265.79
Minosan (Standard)	933.25	1143.15	1174.89	3715.35

13A to 13I, page - 233 to 241  
 [Ref. Tables 14A to 14I, page - 242 to 250]

11. DISCUSSION ON PROTECTANT ACTIVITY AGAINST *H. ORYZAE* ON DETACHED RICE LEAVES.

The protectant activity on detached rice leaves against *H. oryzae* have been carried out by using only two compounds (BD-15 and BD-17). The percent inhibition data have been presented in Table-15.

Table - 15

Code No.	Concentration ( $\mu\text{g/ml}$ )	No. of lesions observed after 48 hours.	Percent inhibition of infection.
BD - 15	1500	22	40.54
	1000	26	29.73
	500	31	16.22
BD - 17	1500	14	62.16
	1000	19	48.65
	500	32	13.51
Control	--	37	--

The results reveal that the nonylamido compound (BD-17) at 1500  $\mu\text{g/ml}$  and 1000  $\mu\text{g/ml}$  inhibited the infection by *H. oryzae* on detached leaves by 62 and 48%, whereas the pyrrolidino compound (BD-15) at the same concentration inhibited the infections by 40 and 29% respectively. The activity in vivo of both the compounds against this pathogen was much lower than that in vitro.

12. DISCUSSION ON PROTECTANT ACTIVITY AGAINST H. ORYZAE ON RICE PLANTS.

The protectant activity on rice plants (40 days old) against *H. oryzae* have been performed by using pyrrolidino (BD-15) and nonylamido (BD-17) compounds. The percent inhibition of ~~infection~~ of infection data have been given in Table-16.

The nonylamido compound (BD-17) at 1500  $\mu\text{g/ml}$  and 1000  $\mu\text{g/ml}$  inhibited the infection by 56% and 44% respectively, whereas the pyrrolidino compound at the same concentrations inhibited the infection by 51 and 47% respectively.

Table - 16

Protectant activity of BD-15 and BD-17 against Helminthosporium oryzae infection on rice plant (40 days old).

Treatment	Concentration ( $\mu\text{g/ml}$ )	Mean no. of spots/ plant <sup>(a)</sup>	Mean disease index/ plant <sup>(a)</sup>	't' at 5% level (P=0.05)	Percent inhibition of infection <sup>(b)</sup>
BD - 15	1500	5.4	2.212	5.87	51.64
	1000	6.0	2.425	5.24	47.04
	500	9.95	3.80	2.13	16.93
BD- 17	1500	4.8	2.0	8.29	56.28
	1000	5.35	2.525	5.39	44.80
	500	7.45	3.10	3.38	32.24
Control	--	12.30	4.575	--	--

(a) = Average of 20 plants.

(b) = Percentage of inhibition of infection =  $\frac{P_n - P_t}{P_n} \times 100$

$P_n$  = Disease index of control plants.

$P_t$  = Disease index of treated plants.

\* = The disease index has been calculated by using the method of Chattopadhyay and Bose [Plant Disease Reporter 63, 103 (1979)]

13. DISCUSSION ON PHYTOTOXIC PROPERTIES

13. A. Seed treatment by soaking.

The toxic effects of the compounds (BR-15 and BR-17) on the germination and root and shoot growth of rice (Dharial variety) are presented in Table - 17.

Treatments of seeds with these two compounds have no effect on germination compared to control. However, root and shoot growth have been drastically reduced for both the compounds. The percent decrease of root length over control is greater than that of shoot length.

13. B. Effect of Spraying on Rice Plants.

The toxic effects of these two compounds on the shoot growth of rice plants are given in Table - 18. The shoot growth has been reduced drastically for both the compounds. The highest decrease in shoot growth is observed in case of nonylamide compound; the decrease is 63.7% over control. In general an increase in the concentration decreases shoot growth of the rice plants.

Table - 17

Effect of treatment with BD-15 and BD-17 on germination and root and shoot growth of rice seeds.

[ By Seed Soaking ]

Code No. of the Compound.	Concentration of the Compound ( $\mu\text{g/ml}$ )	Seed soaked for 2 hours				
		% of germination	Mean root length(mm)	% decrease over control	Mean shoot length (mm)	% decrease over control.
BD - 15	1500	92 (96)	7.8 (11.80)	53.01 (51.84)	7.8 (8.50)	37.60 (51.51)
	1000	93 (95)	10.9 (13.22)	34.30 (46.04)	9.5 (9.44)	24.00 (46.15)
	500	92 (96)	12.5 (13.25)	24.70 (45.92)	10.2 (11.22)	18.40 (35.82)
BD - 17	1500	90 (96)	7.42 (9.26)	55.30 (62.20)	6.5 (7.46)	48.00 (57.44)
	1000	92 (96)	8.75 (13.22)	47.29 (53.96)	8.5 (11.59)	32.00 (33.88)
	500	94 (96)	10.25 (13.54)	33.25 (44.73)	9.48 (11.53)	24.16 (33.94)
Control	--	92 (96)	16.60 (24.50)	- ( - )	12.50 (17.53)	- ( - )

\* Results in parenthesis denote the value of 4 hours.

Table - 13

Phytotoxic activity of BD-15 and BD-17 on rice plant (Oryzae sativa).

[ By Spraying ]

Code name of the Compound	Concentration of the compound ( $\mu\text{g/ml}$ )	Total shoot length (b) (cm)		Length increase (cm)	Mean length increase	% decrease over control.	't' value at 5% level (P = 0.05)
		Before Spray	After Spray				
BD - 15	1500	564.1	582.1	18.0	0.36	55.00	6.58
	1000	551.0	571.8	20.8	0.41	48.75	5.57
	500	570.4	594.4	24.0	0.48	40.00	4.26
BD - 17	1500	508.0	522.9	14.9	0.29	63.75	7.96
	1000	537.6	556.8	19.2	0.38	52.50	6.17
	500	543.1	568.0	24.9	0.49	33.75	4.00
Control	--	533.1	573.3	40.2	0.80	--	--

b = a total of fifty plants.

14. GENERAL CONCLUSIONS AND REMARKS:

(i) Some 2-alkylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphides have been prepared by the reaction of the corresponding phosphoramidethioic dichlorides with 5-nitro-saligenin. All compounds are white crystalline solids. Attempts had also been made several times to prepare the 2-methylamido-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide, and its ethylamido analog; in each case a viscous yellowish liquid was obtained; the pure amidophosphorothionates could not be isolated from these liquid mixtures.

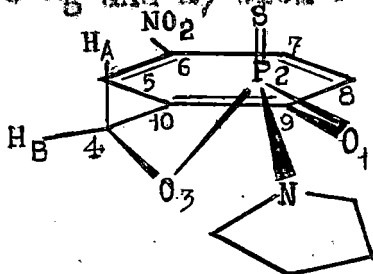
(ii) The structure of the compounds have been determined by chemical analyses and IR, mass and PMR spectra. The spectral data of only four compounds are presented.

The common IR bands for the compounds are : 1010-1030  $\text{cm}^{-1}$ (s), P-O-C (alkyl); 1235-1250  $\text{cm}^{-1}$ (vs), P-O-C (aryl); 880-920  $\text{cm}^{-1}$ (s), P-O-C (aryl); 1515-1520  $\text{cm}^{-1}$ (s), asym. str. of nitro group; 1340-1345  $\text{cm}^{-1}$ (s), sym. str. of nitrogroup; 800-820  $\text{cm}^{-1}$ , P = S (I); 640-660  $\text{cm}^{-1}$ , P = S (II). Neither of the two P = S bands shows any systematic shifts which reflect changes in the inductive properties of the substituent; this is not unexpected if they do indeed arise from mixed modes.

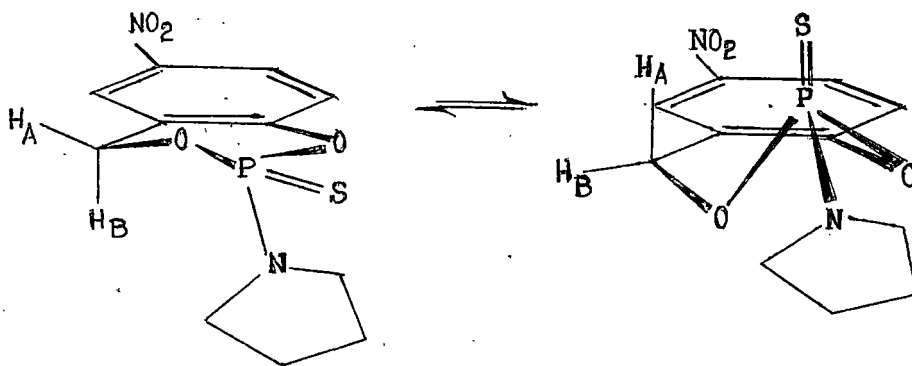
The compounds show parent molecular-ion ( $M^+$ ) peaks in the mass spectra. Fragmentation by loss of 'SH' radical is important; all compounds show an ion due to  $(M - \text{SH})^+$ , and it is the base peak for all of the four alkylamidophosphorothionates.

Finally the structure of the compounds has been settled by taking PMR spectra; for all compounds the endo-cyclic  $-CH_2-$  group gives eight lines in PMR spectra.

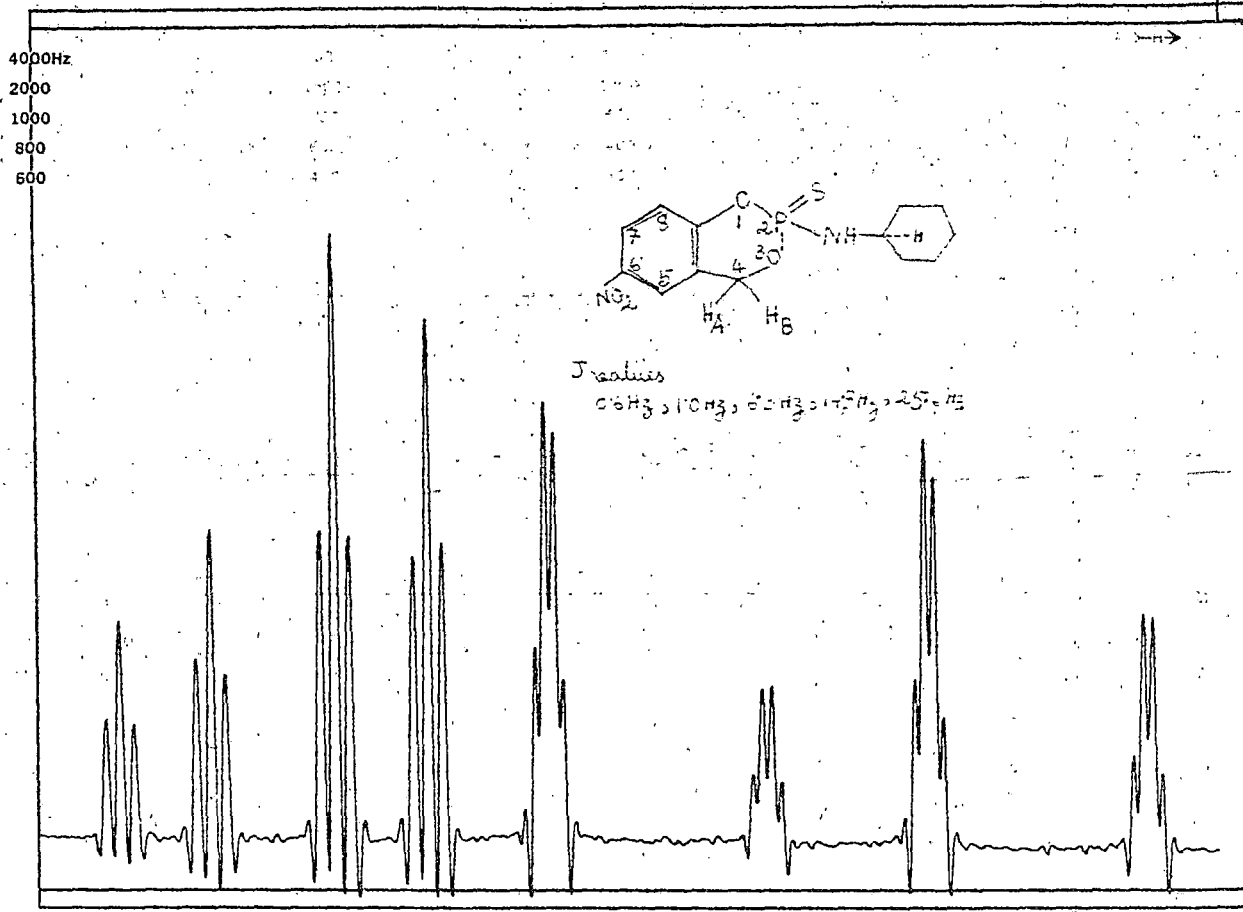
Recently several plot expansions\* and decoupling experiments have been performed in our laboratory (Das et al., unpublished results). These experiments, in case of BD-10, suggest that the quasi-equatorial proton  $H_{4B}$  is assigned to the 5.2 ppm proton ( $J = 25.6 H_z$ ) and the quasi-axial proton  $H_{4A}$  is assigned to the 5.6 ppm. proton ( $J = 6.3 H_z$ ); the geminal coupling constant is  $14.7 H_z$ . It has also been observed that the proton  $H_{4B}$  is coupled equally to the three aromatic protons  $H_5$ ,  $H_7$  and  $H_8$  with  $J = 0.6 H_z$ ; the proton  $H_{4A}$  is more strongly coupled to  $H_5$  and  $H_7$  with  $J = 1.0 H_z$ , but not to  $H_8$



From the temperature dependent PMR spectral study at 270 MHz in the temperature range  $-70^{\circ}C$  to  $+50^{\circ}C$  of the methoxy compound (BD-8), it is fairly evident that the methylene protons of the hetero ring are not equivalent to each other, and the dioxaphosphorinane ring is conformationally mobile in solution (Das et al.,



\* Graphs are attached (Fig- 11, 12, 13).



FT-80A SPECTRUM NO. 14501-1  
 OPERATOR GMS DATE 9/11/79  
 NUCLEUS  $^1\text{H}$  FREQUENCY 125  
 SYNTHESIZER SETTING \_\_\_\_\_  
 EXPERIMENT NAME \_\_\_\_\_  
 FILE NAME BD-10  
 SAMPLE BD-10  
 Plot Expansion of  $\text{CH}_2$  group

LOCK  INTERNAL  EXTER  
 LOCK SIGNAL CQL1  
 SPIN RATE 7.8 105 TEMP 20  
 INSERT 5 mm  $^1\text{H}$

ACQUISITION  
 SPECTRAL WIDTH (SW) \_\_\_\_\_  
 NO. OF TRANSIENTS (NT) \_\_\_\_\_  
 ACQUISITION TIME (AT) \_\_\_\_\_  
 PULSE WIDTH (PW) \_\_\_\_\_  
 PULSE DELAY (PD) \_\_\_\_\_  
 DATA POINTS (DP) \_\_\_\_\_

TRANSMITTER OFFSET (TO) \_\_\_\_\_  
 HIGH FIELD LOW FIELD  
 RECEIVER GAIN (RG) \_\_\_\_\_

DECOUPLER MODE (DM) \_\_\_\_\_  
 DECOUPLER OFFSET (DO) \_\_\_\_\_  
 NOISE BANDWIDTH (NB) \_\_\_\_\_  
 ACQUISITION MODE (AM) \_\_\_\_\_

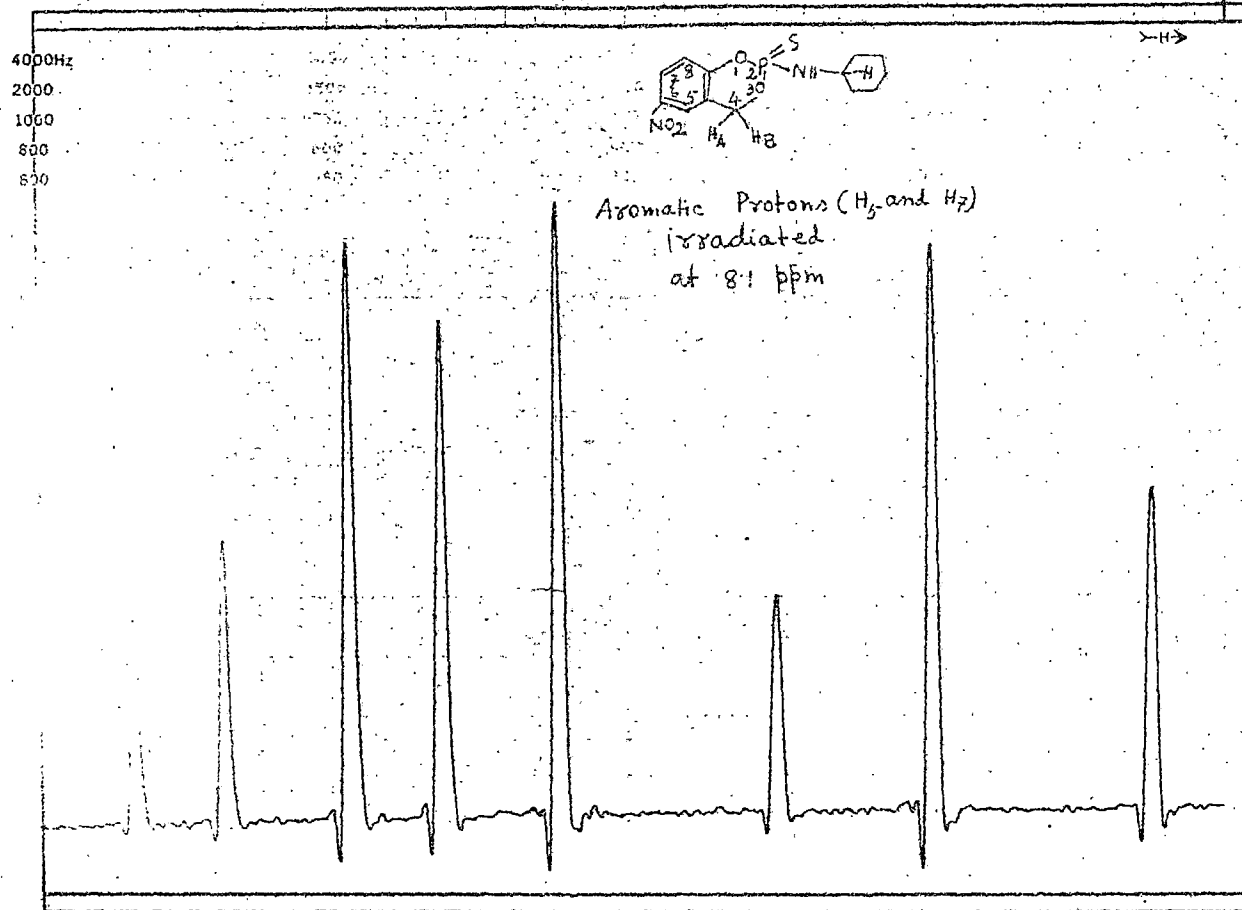
DISPLAY  
 SENS ENHANCEMENT (SE) \_\_\_\_\_  
 WIDTH OF PLOT (WP) 80  
 END OF PLOT (EP) \_\_\_\_\_  
 WIDTH OF CHART (WC) 80  
 END OF CHART (EC) \_\_\_\_\_  
 VERTICAL SCALE (VS) 100  
 REFERENCE LINE (RL) \_\_\_\_\_

Plot Expansion of  $\text{-CH}_2\text{-}$  group in diocaphosphorin ring.

$\text{H}_A \text{H}_B$

Fig-11 Plot expansion of the  $\text{-CH}_2\text{-}$  group (in BD-10),  
 5.0 - 5.8 ppm region.





FT-80A SPECTRUM NO. MSD1-7c  
 OPERATOR \_\_\_\_\_ DATE \_\_\_\_\_  
 NUCLEUS \_\_\_\_\_ FREQUENCY \_\_\_\_\_  
 SYNTHESIZER SETTING \_\_\_\_\_  
 EXPERIMENT NAME \_\_\_\_\_  
 FILE NAME BD-10-A  
 SAMPLE BD-10

Aromatic Protons irradiated  
 $H_5$  and  $H_7$  at 8.1 ppm.

Plot Expansion of  $CH_2$

LOCK  INTERNAL  EXTERNAL  
 LOCK SIGNAL \_\_\_\_\_  
 SPIN RATE \_\_\_\_\_ rps. TEMP. \_\_\_\_\_ °C  
 INSERT \_\_\_\_\_ mm

ACQUISITION  
 SPECTRAL WIDTH (SW) \_\_\_\_\_ Hz  
 NO. OF TRANSIENTS (NT) \_\_\_\_\_  
 ACQUISITION TIME (AT) \_\_\_\_\_ sec.  
 PULSE WIDTH (PW) \_\_\_\_\_ sec.  
 PULSE DELAY (PD) \_\_\_\_\_ sec.  
 DATA POINTS (DP) \_\_\_\_\_

TRANSMITTER OFFSET (TO) \_\_\_\_\_  
 HIGH FIELD \_\_\_\_\_ LOW FIELD \_\_\_\_\_  
 RECEIVER GAIN (RG) \_\_\_\_\_

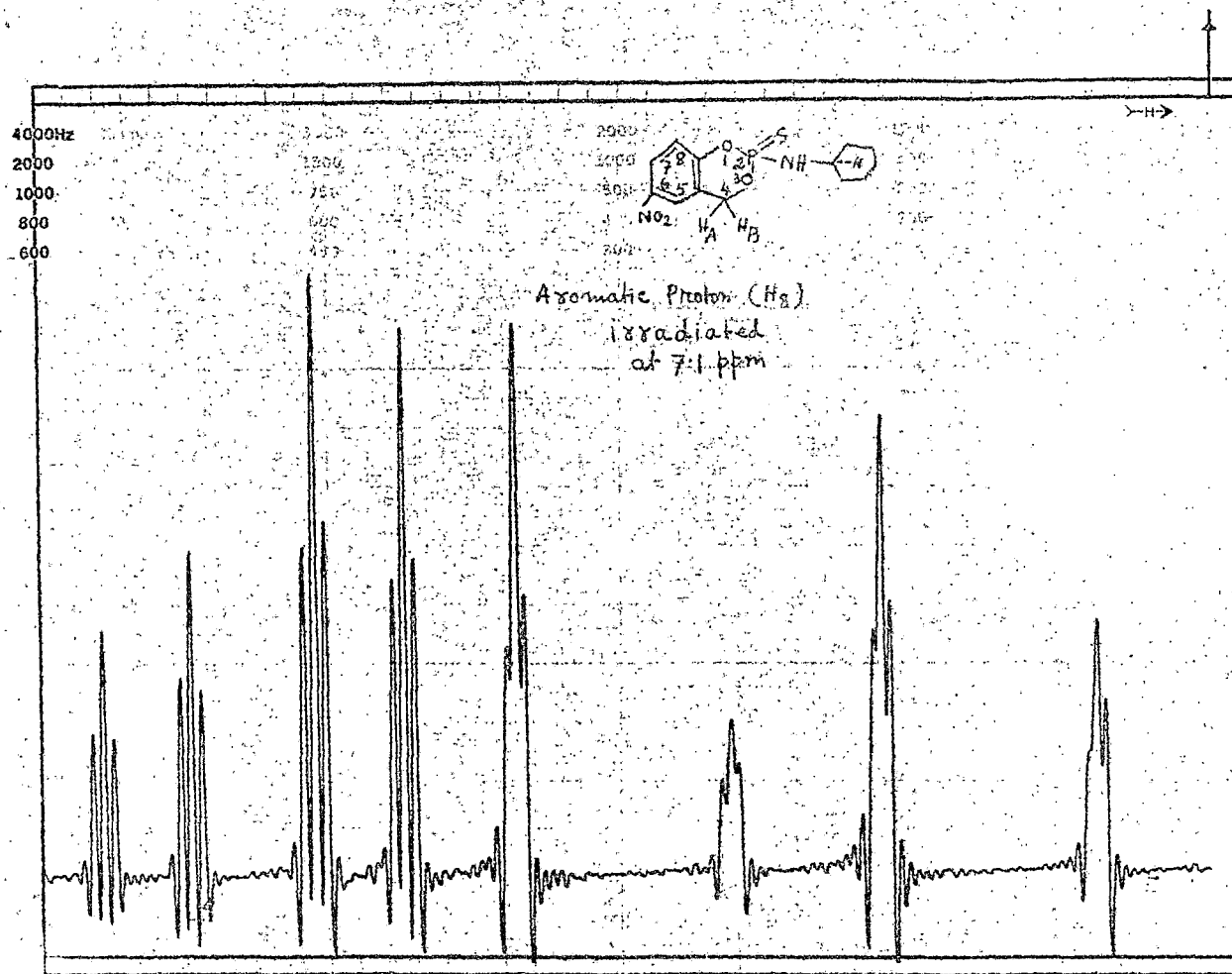
DECOUPLER MODE (DM) \_\_\_\_\_  
 DECOUPLER OFFSET (DO) \_\_\_\_\_  
 NOISE BANDWIDTH (NB) \_\_\_\_\_ kHz  
 ACQUISITION MODE (AM) \_\_\_\_\_

DISPLAY  
 SENS ENHANCEMENT (SE) \_\_\_\_\_ sec.  
 WIDTH OF PLOT (WP) 80 Hz  
 END OF PLOT (EP) \_\_\_\_\_ Hz  
 WIDTH OF CHART (WC) 10 Hz  
 END OF CHART (EC) \_\_\_\_\_ Hz  
 VERTICAL SCALE (VS) 100  
 REFERENCE LINE (RL) \_\_\_\_\_

Plot Expansion of  $CH_2$  group

Fig-12. Plot expansion of  $-CH_2-$  group in BD-10 (aromatic protons  $H_5$  and  $H_7$  irradiated at 8.1 ppm).





FT-60A SPECTRUM NO. 11521-3  
 OPERATOR \_\_\_\_\_ DATE \_\_\_\_\_  
 NUCLEUS \_\_\_\_\_ FREQUENCY \_\_\_\_\_  
 SYNTHESIZER SETTING \_\_\_\_\_  
 EXPERIMENT NAME \_\_\_\_\_  
 FILE NAME BD-10-B  
 SAMPLE BD-10

Aromatic Proton irradiated  
 H<sub>8</sub> at 7.1 ppm  
 Plot expansion of CH<sub>2</sub>

LOCK  INTERNAL  EXTERNAL  
 LOCK SIGNAL \_\_\_\_\_  
 SPIN RATE \_\_\_\_\_ rps. TEMP \_\_\_\_\_ °C  
 INSERT \_\_\_\_\_ mm

ACQUISITION  
 SPECTRAL WIDTH (SW) \_\_\_\_\_ Hz  
 NO. OF TRANSIENTS (NT) \_\_\_\_\_  
 ACQUISITION TIME (AT) \_\_\_\_\_ sec.  
 PULSE WIDTH (PW) \_\_\_\_\_ sec.  
 PULSE DELAY (PD) \_\_\_\_\_ sec.  
 DATA POINTS (DP) \_\_\_\_\_

TRANSMITTER OFFSET (TO):  
 HIGH FIELD \_\_\_\_\_ LOW FIELD \_\_\_\_\_  
 RECEIVER GAIN (RG) \_\_\_\_\_

DECOUPLER MODE (DM) \_\_\_\_\_  
 DECOUPLER OFFSET (DO) \_\_\_\_\_  
 NOISE BANDWIDTH (NB) \_\_\_\_\_ kHz  
 ACQUISITION MODE (AM) \_\_\_\_\_

DISPLAY  
 SENS ENHANCEMENT (SE) \_\_\_\_\_ sec.  
 WIDTH OF PLOT (WP) 80 Hz  
 END OF PLOT (EP) \_\_\_\_\_ Hz  
 WIDTH OF CHART (WC) 80 Hz  
 END OF CHART (EC) \_\_\_\_\_ Hz  
 VERTICAL SCALE (VS) 100  
 REFERENCE LINE (RL) \_\_\_\_\_

Plot Expansion of  $\text{-CH}_2\text{-}$  group

Fig-13. Plot expansion of  $\text{-CH}_2\text{-}$  group in BD-10 (aromatic proton H<sub>8</sub> irradiated at 7.1 ppm).



unpublished results).

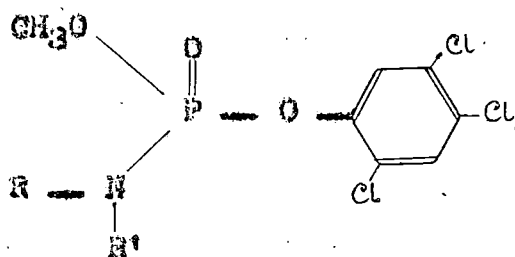
- (iii) The dimethylamido (BD-13) and the morpholine (BD-11) compounds have some insecticidal activity; but their activities are less than that of the 2-methoxy-6-nitro-4H-1,3,2-benzodioxaphosphorin-2-sulphide (BD-8) and Salithion. The other compounds are non-insecticidal.
- (iv) All compounds (BD-10 to BD-15) are less toxic to rats than salithion; however, the diethylamido (BD-12), the dimethylamido (BD-13), and the pyrrolidino (BD-15) compounds have greater toxicity compared to other phosphoramidothionates.
- (v) The results of wheat seed germination studies indicate that none of the compounds are phytotoxic upto 500 ppm concentration, however some of the compounds (BD-10, BD-11 and BD-17) show 90 percent germination at 500 ppm.

Treatments of rice seeds with the compounds (BD-15 and BD-17) have no effect on germination; however, root and shoot growth have been drastically reduced. Further studies on the toxic effects of these two compounds (BD-15 and BD-17) on rice plants indicate that the shoot growth has been reduced drastically for both the compounds. Phytotoxic properties of other compounds on rice seeds and plants have not yet been performed.

- (vi) It has been observed that for any compounds (BD-10 to BD-14), the housefly-head acetylcholinesterase (HFAChE) is more inhibited than the blood-cholinesterase (blood-ChE). The antiacetylcholinesterase

activity of the dimethylamidophosphorothionate (BD-13) is highest, and that of the diethylamidophosphorothionate (BD-12) is least (for AChE); but the anticholinesterase activity for blood - ChE of the morpholino compound (BD-11) is most, and that of the isopropylamidophosphorothionate (BD-14) is least.

Hansch and Deutsch (Biochim. Biophys. Acta., 121, 117, 1966) analysed the data obtained by Fukuto et al. (J. Econ. Entomol., 51, 808, 1963) from a series of methyl 2,4,5-trichlorophenyl - N - alkyl phosphoramidates in order to clarify the effect of the N - alkyl group on the inhibitory activity of the AChE. They observed that



the logarithm of the bimolecular inhibition constant is correlated excellently with Taft's steric constant  $H_s$  and polar constant  $\sigma^*$  of the substituent; the bulky isopropyl and tert - butyl groups decrease inhibition rates by steric interference. On the other hand the ring substituents of methyl phenyl N-methyl phosphoramidates directly affect the antiacetylcholinesterase activity (AChE) by virtue of the electronic and hydrophobic properties (Neely et al, J. Agric. Food Chem., 16, 571, 1968). However, in a series of ethyl S-(substituted)-phenyl phosphoramidates, no correlation was observed between the rates of cholinesterase inhibition

dition and any of the free energy parameters for ring substituents (Sanborn, J.R. and Fucato, T.R; J. Agric. Food. Chem., 20 926, 1972); moreover, the anticholinesterase activity (H<sup>+</sup>AChE) of phosphoramidothiolates is not always correlated with their insecticidal activity.

In case of nitro-saligenin cyclic phosphoramidothionates, the antiacetylcholinesterase activity (H<sup>+</sup>AChE) is not correlated with their insecticidal activity. Among the eight compounds (BD-10 to BD-18), only the dimethylamido compound (BD-13) shows highest insecticidal activity, and also highest antiacetylcholinesterase activity. Although antiacetylcholinesterase activity of both dimethylamido and isopropylamido compounds are comparable ( $I_{50}$  is  $1.80 \times 10^{-5}$  M and  $1.86 \times 10^{-5}$  M respectively, Table - 6, column - 2, page-138), the insecticidal activity of the dimethylamido compound is highest ( $LC_{100} = 3.5 \mu\text{g/g}$ ) and that of the isopropylamido compound is least ( $LC_{100} = >10 \mu\text{g/g}$ ). When the data for other nitro-saligenin cyclic phosphoramidothionates will be available we will try to find out the correlation between antiacetylcholinesterase activity and  $E_s$ ,  $\sigma^+$  as well as  $\pi$  values of alkylamido groups.

(vii) From the chemical hydrolysis studies it has been observed that the compounds containing the disubstituted amido groups are extremely resistant to hydrolysis compared to other compounds having the monosubstituted amido groups. It has also been observed that the rate of alkaline hydrolysis is increased as the pH value increases from 7.7 to 11.8.

(viii) From the fungicidal activity studies (by growth inhibition) against F. oryzae, V. albo-atrum, A. solani and H. oryzae indicate that some of the compounds show good inhibitory effect on the growth of different fungi; however, compared to Hinosan they have less inhibitory effect. Among the nine compounds, the dimethylamido compound is most active against F. oryzae, the isopropylamido compound is against H. oryzae, and the cyclohexylamido compound is against V. albo-atrum & A. solani.

From the spore germination inhibition studies against A. niger, F. gensei, V. albo-atrum and H. oryzae, it has been observed that all compounds are effective. The nonylamido compound is most active for A. niger, and the dimethylamido compound is against F. gensei & V. albo-atrum; their activities are greater than that of Hinosan. The isopropylamido compound is most effective against H. oryzae; but, its activity is less than that of Hinosan.

Protectant activity studies ( in vivo ) against H. oryzae on detached rice leaves and rice plants by using only two compounds (BD-15 and BD-17) indicate that the activity of nonylamido compound (BD-17) is greater than that of the pyrrolidino compound (BD-15). Protectant activity studies for other compounds have not yet been performed.

(ix) The antifungal activity data justify further examination of these phosphoramidothionates as potential fungicides with special reference to the selectivity of their action. Whether the use of these compounds will protect the plants from diseases in the field remains to be studied.

In order to find out the chemical structure — biological activity relationship in these nitro-saligenin cyclic phosphoramidothionates, we have to synthesize several new compounds in which the nitro group is to be incorporated in different position of the aromatic ring, and to investigate their biological activities, phytotoxic properties, fungicidal activities, anti - SII enzyme activities, and other toxicological properties including delayed neurotoxicity in hens.

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T A B L E S

Table - 1A

Effect of BD - 10 on the growth of *Pyricularia oryzae*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	88.23*	77.55*	72.60*
150	100	79.41*	69.38*	65.75*
100	100	76.47*	63.33*	59.49*
50	92.50*	60.29*	50.65*	42.75*
25	64.00*	46.45*	36.75*	30.50*
12.5	50.00*	30.75*	24.45*	16.45*
5	25.50*	19.60	11.25	8.75
ED <sub>95</sub> =	60.25	275.42	537.03	616.59
ED <sub>50</sub> =	12.30	30.19	47.86	66.06
Regression constants: $Y = mx + c$				
m =	64.840	46.486	43.104	46.145
c =	- 20.780	- 13.836	- 22.745	- 34.074
r =	0.998	0.997	0.999	0.999

\* Data used for regression analysis.

Table - 1B

Effect of BD - 11 on the growth of Pyricularia oryzae

Concentration ( $\mu\text{g}/\text{ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	90.47*	78.61*
150	100	85.29*	82.85*	70.07*
100	100	76.47*	71.42*	61.66*
50	73.91*	58.75*	52.38*	45.92*
25	56.82*	44.17*	38.09*	26.61*
12.5	43.47*	29.41*	23.50*	19.35*
5	24.50*	17.35	12.45	8.70
ED <sub>95</sub> =	138.03	229.03	251.13	389.04
ED <sub>50</sub> =	16.59	30.90	39.90	61.65
Regression constants: $Y = mx + c$				
m =	48.866	51.955	55.717	56.725
c =	- 9.698	- 27.844	- 39.020	- 51.970
r =	0.998	0.999	0.997	0.998

\* Data used for regression analysis.

Table - 1C

Effect of BD - 12 on the growth of Pyricularia oryzae

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	100
150	100	100	94.36*	87.60*
100	100	100	84.09*	76.92*
50	88.33*	70.58*	64.90*	58.15*
25	70.91*	56.70*	50.27*	40.50*
12.5	51.50*	36.75*	30.50*	20.75*
5	32.33*	17.64*	9.00	7.65
ED <sub>95</sub> =	69.18	134.39	151.35	194.93
ED <sub>50</sub> =	10.47	19.49	25.11	35.48
Regression constants: $Y = mx + c$				
m =	54.894	54.051	57.622	61.414
c =	- 6.319	- 20.255	- 31.136	- 45.654
r =	0.998	0.997	0.999	0.999

\* Data used for regression analysis

Table 13

Effect of ED - 13 on the growth of *Pyricularia oryzae*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	100
150	100	100	100	100
100	100	100	91.83*	91.83*
50	100	100	93.80*	70.65*
25	100	93.76*	71.50*	47.50*
12.5	92.50	71.23*	60.00*	26.75*
6	68.25	44.30*	22.50*	21.50*
ED <sub>96</sub> =	> 12.50	25.70	48.97	134.89
	< 25.00			
ED <sub>50</sub> =	< 6.00	5.88	10.96	21.87
Regression constants: $Y = mx + c$				
m =		70.477	68.958	56.497
c =		- 4.707	- 21.835	- 25.868
r =		0.999	0.988	0.971

\*Data used for regression analysis

Table - 11

Effect of BD - 14 on the growth of *Fusicularia oryzae*.

Concentration ( $\mu\text{g}/\text{ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	100
150	100	100	100	100
100	100	100	100	100
50	100	100	100	89.13
25	100	63.63	52.77	43.47
12.5	100	42.00	32.50	17.39
5	53.60	18.18	16.66	-
$\text{ED}_{95} =$	> 5.00 < 12.50	> 25.00 < 50.00	> 25.00 < 50.00	> 50.00 < 100.00
$\text{ED}_{50}$	< 5.00	> 12.50 < 25.00	> 12.5 < 25.00	> 25.00 < 50.00

Table - 1F

Effect of BD - 15 on the growth of *Pyricularia oryzae*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	100
150	100	100	100	92.39*
100	100	100	96.50*	78.90*
50	90.50*	80.01*	76.45*	60.65*
25	71.25*	62.45*	58.35*	42.85*
12.5	51.42*	46.05*	44.50*	25.35*
5	30.25*	24.25*	13.15*	6.30
ED <sub>95</sub> =	56.83	91.20	93.32	169.82
ED <sub>50</sub>	10.71	14.12	17.87	31.62
Regression constants: $Y = mx + c$				
m =	60.543	55.609	61.941	61.461
c =	-12.705	-14.375	-27.892	-41.990
r =	0.998	0.989	0.996	0.998

\*Data used for regression analysis

Table - 10

Effect of ED - 16 on the growth of Pyricularia oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	86.50*	80.25
150	100	79.75*	78.58*	71.50*
100	100	72.72*	67.72*	60.06*
50	66.66*	54.54*	52.83*	48.00*
25	52.38*	40.45*	34.73*	24.37*
12.5	33.33*	29.30*	15.50*	11.63
5	12.50*	9.75	6.25	4.60
ED <sub>95</sub> =	154.88	275.42	281.83	363.07
ED <sub>50</sub> =	23.44	38.01	45.70	61.65
Regression constants: $Y = mx + c$				
m =	54.979	52.533	57.603	58.654
c =	- 25.592	- 33.473	- 46.183	- 55.193
r =	0.988	0.999	0.999	0.993

\* Data used for regression analysis

Table - III

Effect of BD - 17 on the growth of Pyricularia oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	95.23*
150	100	100	100	86.59*
100	100	88.21*	84.75*	75.30*
50	88.23*	73.68*	65.15*	56.15*
25	70.53*	55.28*	47.25*	36.35*
12.5	50.50*	45.80*	34.50*	14.25*
5	29.41*	15.05*	11.25*	8.35
ED 95 =	63.09	123.02	153.48	194.98
ED50 =	11.22	18.19	25.11	40.73
Regression constants: $Y = mx + c$				
$m =$	59.375	54.166	55.689	66.113
$c =$	-12.461	-12.512	-27.989	-56.611
$r =$	0.998	0.998	0.996	0.999

\* Data used for regression analysis

Table - I I

Effect of BD-18 on the growth of Pyricularia oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	100
150	100	100	95.45*	90.90*
100	100	92.00*	83.30*	77.75*
50	96.42*	75.00*	65.75*	56.23*
25	78.57*	57.50*	46.63*	36.69*
12.5	57.90*	40.00*	27.54*	16.25*
5	32.00*	16.50*	10.20	6.35
ED <sub>95</sub> =	45.70	139.03	147.91	177.82
ED <sub>50</sub> =	9.12	11.22	27.54	38.01
Regression constants: $Y = mx + c$				
m =	64.861	33.124	62.135	68.00
c =	- 12.584	- 9.892	- 39.091	- 53.040
r	0.999	0.913	0.999	0.999

\* Data used for regression analysis

Table - II

Effect of Minosan on the growth of Pyricularia oryzae

Concentration ( $\mu\text{g}/\text{ml}$ )	Percentage of inhibition over control after			
	48 hours	72 hours	96 hours	120 hours
200	100	100	100	100
150	100	100	100	100
100	100	100	100	100
50	100	100	100	100
25	100	100	91.50	87.50
12.8	100	91.73	82.83	78.75
5	84.61	76.85	68.57	62.50
$\text{ED}_{95} =$	$> 5.00$	$> 12.50$	$> 25.00$	$> 25.00$
	$< 18.50$	$< 25.00$	$< 50.00$	$< 50.00$
$\text{ED}_{50} =$	$< 5.00$	$< 5.00$	$< 5.00$	$< 5.00$

Table - 6A

Acetylcholinesterase Inhibition in House-fly head homogenate (HFAChE)  
at 30°C of BD - 10

(Phosphate buffer, pH = 8.0; total volume = 5.15 ml/5 fly head,  
 $\lambda = 625 \text{ nm}$ ; incubation time 30 min.).

Sets	Inhibition* Conc. ( $\mu\text{g}$ )	O. D	$\Delta$ O.D.	% Inhibition (Y)	$I_{50}$ (M)
Control	-	0.57	0	-	
1	60	0.26	0.32	56.13	
2	50	0.28	0.29	50.87	
3	40	0.32	0.25	43.85	$2.8611 \times 10^{-5}$
4	30	0.35	0.22	39.60	
5	20	0.405	0.165	28.90	

X\* = log (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :

$$Y = mx + c$$

$$c = -44.5645$$

$$m = 56.1474$$

$$r = 0.9963$$

Table - 6B

Acetylcholinesterase Inhibition in House-fly head homogenate (HPAChE)  
at 30°C of BA-11

(Phosphate buffer; pH = 8.0; total volume = 5.15 ml/5 fly head,  
 $\lambda = 625 \text{ nm}$ ; in-cubation time 30 min).

Sets	Inhibition* Conc. ( $\mu\text{g}$ )	O. D.	$\Delta$ O. D.	% Inhibition (Y)	$I_{50}$ (M)
Control	(-)	0.610	0	-	
1	50	0.310	0.300	39.21	
2	40	0.340	0.170	33.35	
3	30	0.370	0.140	27.45	$5.6326 \times 10^{-5}$
4	20	0.415	0.095	18.60	
5	10	0.480	0.030	5.88	

$X^* = \log$  (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :

$$Y = mx + c$$

$$c = -42.0753$$

$$m = 47.3029$$

$$r = 0.9983$$

Table - 6C

Acetylcholinesterase Inhibition in House-fly head homogenate (HFACHH)  
at 30°C of 50 - 12.

(Phosphate buffer; pH = 8.0; total volume 5.15 ml/5 fly head,  
 $\lambda = 625 \text{ nm}$ ; in-cubation time 30 min).

Sets	Inhibition* Conc. ( $\mu\text{g}$ )	O.D.	$\Delta$ O.D.	%Inhibition (Y)	$I_{50}$ (10)
Control	(-)	1.05	0	-	
1	100	0.57	0.48	45.71	
2	80	0.59	0.46	43.81	
3	60	0.64	0.41	39.09	
4	40	0.71	0.34	32.38	$8.2601 \times 10^{-5}$
5	20	0.80	0.25	23.81	
6	10	0.92	0.13	12.38	

X\* = log (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :

$$Y = mx + c$$

$$c = -20.6648$$

$$m = 33.5095$$

$$r = 0.9985$$

Table - 6D

Acetylcholinesterase Inhibition in House-fly head homogenate (HPAChE)  
at 30°C of DD-13.

(Phosphate buffer; pH = 8.0; total volume 5.15 ml/5 fly head,  
 $\lambda = 625 \text{ nm}$ ; in-cubation time 30 min).

Sets	Inhibition* Conc. ( $\mu\text{g}$ )	O.D.	$\Delta$ O.D.	Inhibition (Y)	$I_{50}$ (M)
Control	(-)	1.05	0	-	
1	100	0.21	0.84	80.00	
2	80	0.27	0.78	74.28	
3	60	0.33	0.72	68.57	$1.80 \times 10^{-5}$
4	40	0.42	0.63	60.00	
5	20	0.53	0.47	44.76	

$X^* = \log (\text{Conc. of the inhibitor in } \mu\text{g}).$

Regression constants :  $Y = mx + c$   
 $c = -20.6339$   
 $m = 49.8322$   
 $r = 0.9997$

Table - 6B

Acetylcholinesterase Inhibition in House-fly head homogenate (HFACHE)  
at 30°C of BD-14.

(Phosphate buffer, pH = 8.0; total volume 5.15 ml/5 fly head;

$\lambda = 625 \text{ nm}$ ; in-cubation time 30 min.)

Sets	Inhibition* Conc. ( $\mu\text{g}$ )	O.D.	$\Delta$ O.D.	%Inhibition (%)	$I_{50}$ (M)
Control	(-)	1.05	0	..	
1	50	0.41	0.64	60.95	
2	40	0.46	0.59	56.18	
3	30	0.61	0.54	51.42	$4.8642 \times 10^{-5}$
4	20	0.59	0.46	43.81	
5	10	0.71	0.34	32.38	

X\* = log (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :  
 $Y = mx + c$   
 $c = .8.5553$   
 $m = 40.6143$   
 $r = 0.9993$

Table - 6F

Acetylcholinesterase Inhibition in goat-whole blood of BD-10 at 30°C

(Phosphate buffer, pH = 8.0; total volume = 5.15 ml/0.2 ml blood;  
 $\lambda = 625 \text{ nm}$ ; in cubation time 30 min.).

Sets	Inhibitor* Conc. ( $\mu\text{g}$ )	O.D.	$\Delta\text{O.D.}$	% Inhibition (Y)	$I_{50}$ (M)
Control	(-)	0.59	-		
1	30	0.63	0.06	8.69	
2	40	0.60	0.09	13.04	
3	60	0.56	0.13	13.84	$1.810 \times 10^{-4}$
4	80	0.516	0.175	25.56	
5	100	0.48	0.21	30.43	
6	120	0.46	0.23	33.33	

X\* = log (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :  $Y = mX + c$   
 $c = -53.8102$   
 $m = 41.7679$   
 $r = 0.9759$

Table - 6B

Acetylcholinesterase Inhibition in goat whole blood of BD-11 at 30°C.

(Phosphate buffer; pH = 8.0; total volume = 5.15 ml/0.2 ml blood;  
 $\lambda = 625 \text{ nm}$ ; in cubation time 30 min.)

Sets	Inhibitor* Conc. ( $\mu\text{g}$ )	O. D.	$\Delta\text{O.D.}$	% Inhibition (Y)	$I_{50}$ (M)
Control	(-)	0.80	-		
1	30	0.69	0.11	13.75	
2	40	0.65	0.15	18.75	
3	60	0.58	0.22	27.50	$1.0426 \times 10^{-4}$
4	80	0.54	0.26	32.50	
5	100	0.49	0.31	38.75	
6	120	0.45	0.35	43.75	

X\* = log (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :  $Y = mx + c$   
 $c = -59.8146$   
 $m = 49.2530$   
 $r = 0.9968$

Table - 6H

Acetylcholinesterase Inhibition in goat whole blood of BD.12 at 30°C.

(Phosphate buffer, pH = 8.0; total volume = 5.15 ml/0.2 ml blood;  
 $\lambda = 625 \text{ nm}$ ; in-cubation time 30° min.).

Sets	Inhibitor* Conc. ( $\mu\text{g}$ )	O. D.	$\Delta$ O.D.	% Inhibition (Y)	$I_{50}$ (M)
Control	(-)	0.65	(-)		
1	40	0.59	0.06	9.23	$1.4259 \times 10^{-4}$
2	60	0.53	0.12	18.46	
3	80	0.48	0.17	26.15	
4	100	0.45	0.20	30.76	
5	120	0.42	0.23	35.38	

$X^* = \log$  (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :  $Y = mx + c$   
 $c = -78.8637$   
 $m = 54.9211$   
 $r = 0.9995$

Table - 61

Acetylcholinesterase Inhibition in goat-whole blood of BD-13 at 30°C.

(Phosphate buffer, pH = 8.0; total volume = 5.15 ml/0.2 ml blood,  
 $\lambda = 625 \text{ nm}$ ; in-cubation time 30 min.).

Sets	Inhibition* Conc. ( $\mu\text{g}$ )	O. D.	$\Delta\text{O.D.}$	% Inhibition (Y)	$I_{50}$ (M)
Control	(-)	0.69	-		
1	30	0.65	0.04	6.15	
2	40	0.62	0.07	10.76	
3	60	0.58	0.11	15.94	
4	80	0.55	0.14	20.28	$4.6564 \times 10^{-4}$
5	100	0.53	0.16	23.18	
6	120	0.51	0.18	26.08	

$X^* = \log$  (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :  $Y = mx + c$

$c = -41.7889$

$m = 32.5769$

$r = 0.9996$

Table - 6J

Acetylcholinesterase Inhibition in goat-whole blood of BD-14 at 30°C

(Phosphate buffer, pH = 8.0; total volume = 5.15 ml/0.2 ml blood;  
 $\lambda = 625 \text{ nm}$ ; in cubation time 30 min. ).

Sets	Inhibitor* Conc. ( $\mu\text{g}$ )	O.D.	$\Delta$ O.D.	% Inhibition (Y)	$I_{50}$ (M)
Control	(-)	0.55	-		
1	50	0.47	0.08	14.54	
2	100	0.42	0.13	23.63	
3	150	0.39	0.16	29.09	$4.7805 \times 10^{-4}$
4	200	0.37	0.18	32.72	
5	250	0.33	0.20	36.36	

$X^* = \log$  (Conc. of the inhibitor in  $\mu\text{g}$ ).

Regression constants :  $Y = mx + c$   
 $c = -33.0805$   
 $m = 30.8911$   
 $r = 0.9986$

Table - 7A

Chemical hydrolysis of DL-11 at  $\lambda = 410$  nm; pH = 11.85; 0.0095 M NaOH (in 50% Ethanol)  $\epsilon = 20090$ , Initial concentration,  $C_0 = 7.06 \times 10^{-5}$  M. temperature 20°C.

Time (hrs) (X)	O. D.	$C_t \times 10^5$ (M)	$\log \frac{C_0}{C_0 - C_t}$ (Y)	$K_{hyd}$ (min <sup>-1</sup> )
5	0.103	0.51269	0.0327	6.1413 x 10 <sup>-5</sup>
27.75	0.316	1.5729	0.1085	
52.50	0.445	2.2150	0.1635	
75.33	0.49	2.4390	0.1841	
124.6	0.6	2.9866	0.2388	

Regression constants :

$$Y = mx + c$$

$$c = 0.0526$$

$$m = 0.0016$$

$$r = 0.9599$$

Table - 7B

Chemical hydrolysis of BD - 13 at  $\lambda = 410$  nm, pH = 11.85, 0.0095 M NaOH (in 50% Ethanol)  $\epsilon = 20000$ , Initial Concentration,  $C_0 = 7.3 \times 10^{-5}$ , temperature  $20^\circ\text{C}$ .

Time (hrs) (X)	O. D.	$C_t \times 10^5$ (M)	$\log \frac{C_0}{C_0 - C_t}$ (Y)	$K_{\text{hyd}}$ ( $\text{min}^{-1}$ )
5	0.034	0.16924	0.0102	
28	0.143	0.71180	0.0446	
52.75	0.210	1.0453	0.0671	$2.687 \times 10^{-5}$
75.25	0.255	1.2693	0.0830	
124.75	0.305	1.5182	0.1013	

Regression constants :  $Y = mx + c$

$c = 0.0103$

$m = 0.0007$

$r = 0.9559$

Table - 7C

Chemical hydrolysis of BD - 15 at  $\lambda = 410$  nm, pH = 11.85, 0.0095 M NaOH (in 50% Ethanol), Initial concentration,  $C_0 = 2.12 \times 10^{-5}$  M, temperature  $20^\circ\text{C}$ .

Time (hrs) (X)	O. D.	$C_t \times 10^5$	$\log \frac{C_0}{C_0 - C_t}$ (Y)	$K_{\text{hyd}}$ ( $\text{min}^{-1}$ )
150	0.063	0.31359	0.0695	$1.7784 \times 10^{-5}$

\* This compound (BD-15) was extremely resistant to hydrolysis at pH 11.85; only after 150 hours reading was taken at  $\lambda = 410$  nm and  $K_{\text{hyd}}$  was calculated from the reading with the help of 1st order rate equation.

$$K = \frac{1}{t} \ln \frac{C_0}{C_0 - C_t}$$

Table - 7D

Chemical hydrolysis of BD-16 at  $\lambda = 410$ , pH = 11.85, 0.0095 M NaOH (in 50% Ethanol), Initial concentration,  $C_0 = 1.67 \times 10^{-6}$  M, temperature  $20^\circ\text{C}$ .

Time (hrs) (X)	O. D.	$C_t \times 10^5$ (M)	$\log \frac{C_0}{C_0 - C_t}$ (Y)	$K_{\text{hyd}}$ ( $\text{min}^{-1}$ )
150	0.044	0.21901	0.0474	$1.2129 \times 10^{-5}$

\* This compound (BD-16) also is extremely resistant to hydrolysis at pH 11.85; only after 150 hours reading was taken at  $\lambda = 410$  nm and  $K_{\text{hyd}}$  was calculated from the reading with the help of 1st order rate equation:

$$K = \frac{1}{t} \ln \frac{C_0}{C_0 - C_t}$$

Table - 7E

Chemical hydrolysis of BA-17 at  $\lambda = 410 \text{ nm}$ ,  $\text{pH} = 11.85$ ,  $0.0096 \text{ M}$   
NaOH (in 50 50% Ethanol)  $\epsilon = 20090$ , Initial concentration,  
 $C_0 = 3.14 \times 10^{-6} \text{ M}$ , temperature  $20^\circ\text{C}$ .

Time (min) (X)	O.D.	$C_t \times 10^5$ (M)	$\log \frac{C_0}{C_0 - C_t}$ (Y)	$K_{\text{hyd}}$ ( $\text{min}^{-1}$ )
5.25	0.07	0.34843	0.0511	
6	0.077	0.33323	0.0566	
10	0.127	0.63216	0.0976	
15	0.177	0.88194	0.1430	
20	0.227	1.085	0.1852	$20.266 \times 10^{-3}$
25	0.265	1.3191	0.2362	
30	0.299	1.4783	0.2764	
35	0.333	1.6575	0.3259	
40	0.358	1.7670	0.3593	
45	0.378	1.8815	0.3971	

Regression constants:

$$Y = mx + c$$

$$c = 0.0083$$

$$m = 0.0088$$

$$r = 0.9911$$

Table - 8A

Effect of BD - 10 on growth of Verticillium- albo - atrum

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	93.05	91.37*
400	100	88.54*	83.05*
300	90.62*	75.35*	70.45*
250	85.41*	69.50*	66.50*
200	81.25*	62.35*	58.75*
150	66.50*	51.50*	48.65*
100	52.08*	39.60*	34.40*
50	27.00*	19.82	12.30
ED <sub>95</sub> =	316.22	501.18	549.54
ED <sub>50</sub> =	91.20	138.03	154.88
Regression constants: $Y = mx + c$			
m =	83.755	79.718	81.187
c =	- 114.606	- 120.719	- 128.023
r =	0.997	0.998	0.998

\* Data used for regression analysis

Table - 8BEffect of ED - 11 on growth of Verticillium albo-atrum.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	100	88.27*
400	100	90.35*	79.50*
300	100	77.75*	66.35*
250	100	69.25*	58.75*
200	90.75*	61.35*	48.90*
150	80.00*	49.90*	41.50*
100	62.25*	37.50*	24.75*
50	34.00*	21.65	9.75
ED <sub>95</sub>	213.79	457.08	538.84
ED <sub>50</sub>	70.79	141.25	190.54
Regression constants: $Y = mx + c$			
m =	93.403	88.507	91.292
c =	- 123.053	- 141.133	- 158.464
r =	0.993	0.997	0.997

\* Data used for regression analysis.

Table - 8C

Effect of BD - 12 on growth of Verticillium albo-atrum.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	87.90*	78.70*
400	93.75*	78.60*	70.83*
300	82.80*	67.15*	60.30*
250	70.31*	57.30*	50.16*
200	63.50*	51.10*	44.50*
150	50.25*	40.70*	30.16*
100	32.81*	23.50*	21.36
50	24.90	17.65	6.66
ED <sub>95</sub> =	398.10	588.84	707.94
ED <sub>50</sub> =	144.54	180.54	234.42
Regression constants: $Y = mx + c$			
m =	102.254	91.872	92.702
c =	- 171.850	- 160.096	- 170.110
r =	0.998	0.998	0.997

\* Data used for regression analysis.

Table - 8D

Effect of BD - 13 on growth of Verticillium albo-atrum.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	100	87.93*
400	100	87.65*	80.01*
300	92.93*	76.50*	68.40*
250	82.50*	68.40*	59.90*
200	75.22*	60.75*	53.50*
150	64.50*	48.35*	40.50*
100	43.24*	32.25*	26.50*
50	19.50*	11.25	7.35
ED <sub>95</sub> =	323.59	467.73	575.43
ED <sub>50</sub> =	104.71	151.35	181.97
Regression constants: $Y = mx + c$			
m =	91.492	92.582	89.695
c =	- 134.907	- 152.629	- 153.421
r =	0.999	0.999	0.999

\* Data used for regression analysis.

Table - 8E

Effect of BD - 14 on growth of *Verticillium albo-atrum*

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	93.00*	86.43*
400	100	87.77*	79.69*
300	88.33*	75.25*	65.72*
250	80.00*	66.66*	56.79*
200	73.33*	56.25*	49.75*
150	62.00*	46.50*	36.25*
100	49.66*	34.75*	20.15*
50	20.15*	15.50	9.25
ED <sub>95</sub> =	354.81	501.18	598.84
ED <sub>50</sub> =	104.71	154.88	199.52
Regression constants : $Y = mx + c$			
m =	86.077	88.487	85.732
c =	- 124.709	- 144.355	- 173.409
r =	0.999	0.996	0.999
* Data used for Regression analysis.			

Table - 8F

Effect of BD - 15 on growth of *Verticillium albo-atrum*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	90.30*	87.50*
400	100	81.81*	78.04*
300	88.31*	72.72*	69.35*
250	80.39*	64.35*	60.45*
200	73.15*	55.55*	53.50*
150	62.00*	43.25*	39.90*
100	49.75*	28.50*	27.65*
50	30.25*	21.65	13.25
ED <sub>95</sub> =	380.19	537.03	602.55
ED <sub>50</sub> =	95.49	169.82	181.97
Regression constants : $Y = mx + c$			
m =	74.073	90.154	87.379
c =	- 86.764	- 151.693	- 147.079
r =	0.296	0.999	0.998
* Data used for regression analysis.			

Table - 80

Effect of ED - 16 on growth of Verticillium albo-atrum.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	95.06*	89.02*
400	100	88.47*	80.62*
300	100	75.45*	68.65*
250	87.14*	68.60*	60.25*
200	80.00*	58.25*	52.50*
150	66.50*	46.00*	38.15*
100	49.75*	30.75*	22.35*
50	20.00*	16.20	6.25
ED <sub>95</sub> =	238.40	467.73	594.54
ED <sub>50</sub> =	97.72	158.48	190.54
Regression constants : $Y = mx + c$			
m =	96.923	95.133	97.189
c =	- 148.830	- 159.792	- 171.963
r =	0.999	0.999	0.999

\* Data used for regression analysis.

Table - 8H

Effect of BD - 17 on growth of Verticillium albo-atrum.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after		
	24 hours	48 hours	62 hours
500	100	83.83*	83.25*
400	88.11*	79.51*	73.43*
300	78.25*	69.25*	60.25*
250	70.50*	62.75*	50.90*
200	63.25*	54.50*	42.30*
150	52.40*	44.25*	27.84*
100	37.22*	29.12*	16.25
50	27.15	16.50	8.00
ED <sub>95</sub> =	437.73	588.84	630.95
ED <sub>50</sub> =	138.03	173.78	234.42
Regression constants : $Y = mx + c$			
m =	85.070	85.307	106.079
c =	- 132.550	- 141.343	- 202.152
r =	0.999	0.999	0.999
* Data used for regression analysis.			

Table - 31

Effect of  $BD_{50}$  - 13 on growth of Verticillium albo-atrum.

Concentration ( g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	62 hours	96 hours
500	83.88*	80.46*	75.60*	
400	77.77*	72.93*	68.20*	
300	69.41*	63.42*	57.25*	
250	63.74*	56.25*	53.75*	
200	58.25*	50.20*	44.75*	
150	46.25*	40.25*	34.25*	
100	34.25*	27.25*	20.50*	
50	21.15	13.20	7.25	
$ED_{95}$ =	676.08	758.57	851.13	
$BD_{50}$ =	162.12	194.93	229.03	
Regression constants: $Y = mx + c$				
$m =$	72.653	76.753	79.754	
$c =$	- 110.846	- 126.425	- 138.780	
$r =$	0.999	0.999	0.998	
* Data used for regression analysis.				

Table - BJ

Effect of Ninosan on growth of *Verticillium albo-atrum*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	62 hours	86 hours
500	100	100	100	
400	100	100	100	
300	100	94.66*	86.50*	
250	100	89.04*	80.50*	
200	90.45	78.15*	70.25*	
150	71.65	66.25*	57.28*	
100	63.25	50.65*	40.45*	
50	41.30	22.60*	13.50*	
ED <sub>95</sub> =	> 200.00	295.12	346.73	
ED <sub>50</sub> =	> 50.00	95.49	123.02	
Regression constants : $Y = mx + c$				
m =		93.313	99.808	
c =		-135.598	-158.215	
r =		0.999	0.999	

\* Data used for regression analysis.

Table - 9A

Effect of DL-10 on growth of Alternaria solani

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	100	90.50*
400	100	100	90.50*	83.05*
300	100	100	81.75*	72.25*
250	100	100	74.25*	65.25*
200	100	82.05*	66.92*	55.35*
150	82.15	71.79*	56.51*	46.41*
100	71.85	58.97*	44.05*	42.52*
50	41.50	35.00*	23.60	14.79
ED <sub>95</sub> =	> 150.00	288.40	446.68	588.84
ED <sub>50</sub> =	> 50.00	75.85	120.22	147.91
Regression constants : $Y = mx + c$				
m =		76.974	78.755	74.133
c =		- 95.075	- 113.847	- 110.968
r =		0.999	0.999	0.983

\* Data used for regression analysis.

Table - 9B

Effect of BD - 11 on growth of Alternaria solani

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	85.29*	80.43*
400	100	90.42*	77.58*	70.20*
300	100	82.14*	64.50*	57.60*
250	100	72.61*	54.04*	43.91*
200	100	63.50*	43.00*	40.82*
150	71.07	57.90*	36.17*	27.25*
100	53.10	50.00*	20.25*	16.15
50	36.25	37.50*	11.35	8.15
ED <sub>95</sub> =	> 150.00	575.43	616.59	676.08
ED <sub>50</sub> =	< 100.00	147.91	209.92	245.47
Regression constants + $Y = mx + c$				
m =		76.239	95.417	101.306
c =		- 116.148	- 171.444	- 192.649
r =		0.996	0.999	0.999

\* Data used for Regression Analysis.

Effect of DD - 12 on growth of Alternaria solani

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	100	91.50*
400	100	100	90.50*	80.75*
300	100	85.20*	78.75*	68.18*
250	100	77.75*	71.42*	60.36*
200	77.77*	70.37*	62.63*	50.00*
150	68.50*	56.50*	50.25*	36.36*
100	52.05*	40.03*	34.26*	22.50*
50	25.25*	12.36	12.30	7.75
ED <sub>95</sub> =	309.02	353.07	436.51	549.84
ED <sub>50</sub> =	93.32	125.89	144.54	194.98
Regression constants: $Y = mx + c$				
m =	85.933	93.600	93.296	99.919
c =	- 119.786	- 152.925	- 151.868	- 178.859
r =	0.999	0.999	0.999	0.999

\* Data used for regression analysis.

Table 92

Effect of ED - 13 on growth of Alternaria solani

Concentration ( g/mL)	Percentage of inhibition over control after				96 hours
	24 hours	48 hours	72 hours	96 hours	
500	100	100	78.50*	69.50*	
400	100	100	70.50*	61.50*	
300	100	79.72*	59.65*	50.65*	
250	100	72.72*	52.00*	44.55*	
200	100	64.50*	44.25*	36.75*	
150	100	52.50*	32.45*	26.15*	
100	68.50	37.65*	18.50*	12.08	
50	37.15	19.15	10.20	6.25	

ED<sub>95</sub> = > 100.00      426.57      741.31      1000.00  
 ED<sub>50</sub> = > 50.00      134.89      229.09      281.83

Regression constants :  $Y = mx + c$

m = 89.909      87.403      82.992  
 c = - 142.316      - 156.828      - 154.042  
 r = 0.999      0.999      0.999

\* Data used for regression analysis.

Table - 9E

Effect of BD - 15 on growth of *Alternaria solani*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	88.71*	82.53*
400	100	87.40*	81.63*	74.90*
300	100	86.44*	70.55*	65.25*
250	100	80.25*	66.30*	55.60*
200	100	70.27*	58.30*	48.00*
150	100	58.75*	46.25*	34.25*
100	85.00	44.50*	34.25*	20.75*
50	62.35	36.15	18.50	8.30
ED <sub>95</sub> =	> 100.00	363.07	575.43	645.65
ED <sub>50</sub> =	< 50.00	114.81	158.48	208.92
Regression constants : $Y = mx + c$				
m =		89.440	79.595	91.406
c =		-134.716	-125.126	-162.556
r		0.999	0.999	0.998

\* Data used for regression analysis.

Table - 97

Effect of ED - 16 on growth of *Alternaria spiani*

Concentration ( g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	100	88.75*
400	100	100	91.70*	81.25*
300	100	90.00*	80.64*	70.00*
250	100	85.00*	73.80*	64.00*
200	100	77.50*	66.25*	56.15*
150	100	69.50*	55.00*	44.20*
100	80.00	52.50*	40.45*	30.62*
50	56.50	28.00	19.15	11.45
ED <sub>95</sub> =	> 100.00	323.59	426.57	575.43
ED <sub>50</sub> =	< 50.00	89.12	128.82	169.82
Regression constants : $Y = mx + c$				
m =		80.583	85.420	84.645
c =		-107.783	-130.344	-139.894
r =		0.998	0.999	0.999
* Data used for regression analysis.				

Table - 90

Effect of BB - 17 on growth of *Alternaria solani*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	76.66*	71.18*
400	100	100	68.00*	62.25*
300	100	71.05*	55.00*	49.15*
250	100	63.52*	47.50*	42.93*
200	100	55.36*	38.66*	33.89*
150	66.66	42.10*	28.50*	23.50*
100	50.00	26.31*	22.50	16.55
50	26.50	15.10	9.75	5.00
$ED_{95} =$	> 150.00	512.86	776.24	891.25
$ED_{50} =$	100.00	173.78	257.03	288.40
Regression constants : $Y = mx + c$				
$m =$	32.004	95.662	93.840	92.066
$c =$		-165.102	-176.280	-177.176
$r =$		0.999	0.999	0.999
* Data used for regression analysis.				

Table - 9H

Effect of BD - 18 on growth of *Alternaria solani*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	89.13*	77.90*
400	100	92.50*	79.50*	69.50*
300	100	81.42*	67.50*	55.32*
250	100	75.70*	58.50*	47.60*
200	100	66.90*	51.08*	44.61*
150	75.00	55.65*	39.13*	30.76*
100	62.50	44.44*	26.78*	20.00
50	50.00	18.50*	13.50	9.50

ED <sub>95</sub> =	>150.00	245.47	598.84	776.24
ED <sub>50</sub> =	50.00	77.62	186.20	239.88

Regression constants :  $Y = mx + c$

m =	80.981	91.119	89.203
c =	- 118.552	- 157.540	- 163.051
r =	0.999	0.997	0.994

\* Data used for regression analysis.

Table - 91

Effect of Hinosan on growth of Alternaria solani

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	100	95.83*
400	100	100	100	90.62*
300	100	100	80.05*	74.50*
250	100	100	87.50*	68.33*
200	100	100	77.15*	57.08*
150	100	86.81*	63.68*	45.70*
100	80.00	70.83*	48.44*	20.50*
50	60.00	40.15*	17.55*	10.25
ED <sub>95</sub> =	> 100.00	177.82	309.02	446.68
ED <sub>50</sub> =	< 50.00	60.25	104.71	169.82
Regression constants: $Y = mx + c$				
m =		97.423	95.599	108.484
c =		- 124.369	- 143.890	- 192.800
r =		0.999	0.998	0.995

\* Data used for regression analysis.

Table - 10A

Effect of BD - 10 on growth of Helminthosporium oryzae.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	91.50*	87.70*
400	100	100	77.50*	76.50*
300	100	76.74*	67.50*	65.81*
250	100	72.09*	62.50*	58.97
200	100	65.11*	52.50*	50.42*
150	100	56.50*	40.00*	35.04*
100	85.71	41.50*	27.50*	24.78*
50	57.14	18.60*	17.50*	11.11*
ED <sub>95</sub>	> 100.00	500.00	575.43	602.55
ED <sub>50</sub>	< 50.00	125.89	181.97	194.93
Regression constants : $Y = mx + c$				
m =		75.645	90.563	92.127
c =		-109.063	- 155.167	- 161.705
r =		0.999	0.995	0.996
* Data used for regression analysis.				

Table - 10B

Effect of BD - 11 on growth of Helminthosporium oryzae.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	72.41*	63.26*	53.84*
400	100	62.05*	54.03*	52.30*
300	80.76*	53.44*	44.89*	43.07*
250	69.23*	44.82*	40.50*	33.77*
200	62.50*	36.50*	26.53*	26.15*
150	51.50*	27.50*	18.36*	16.92*
100	38.46*	17.24*	14.28*	12.30*
50	26.50*	13.73*	10.20*	8.25*
$ED_{95} =$	562.34	977.23	1122.01	1548.81
$ED_{50} =$	123.02	269.15	338.84	389.04
Regression constants : $Y = mx + c$				
$m =$	68.319	80.328	86.680	74.605
$c =$	- 93.429	- 145.87	- 169.942	-143.279
$r =$	0.978	0.995	0.992	0.930
* Data used for regression analysis.				

Table - 10CEffect of ED - 13 on growth of Helminthosporium oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	80.42*	74.15*	67.25*
400	100	71.35*	66.15*	59.37*
300	100	58.92*	53.19*	47.50*
250	100	53.57*	45.74*	39.00*
200	100	45.50*	37.23*	30.25*
150	54.85	33.50*	27.65*	20.15*
100	45.50	18.50*	12.85*	9.25
50	36.15	11.25	8.07	3.15
ED <sub>95</sub> =	>150	724.43	851.13	954.99
ED <sub>50</sub> =	>100	293.87	263.02	316.22
Regression constants: $Y = mx + c$				
m =		88.726	88.447	92.368
c =		- 159.028	-164.848	- 181.150
r =		0.999	0.999	0.999

\* Data used for regression analysis.

Table - 100

Effect of BD - 14 on growth of Helminthosporium oryzae.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	100	100
400	100	100	92.00*	81.50*
300	100	100	82.15*	60.50*
250	100	92.50*	77.50*	59.65*
200	100	82.25*	65.50*	51.50*
150	100	74.15*	53.75*	42.65*
100	68.25	56.20*	38.15*	28.75*
50	53.10	28.50*	16.05	7.80
ED <sub>95</sub> =	> 100.00	263.02	407.33	620.95
ED <sub>50</sub> =	< 50.00	83.17	131.82	186.20
Regression constants : $Y = mx + c$				
m =		90.755	91.937	84.581
c =		- 124.770	- 145.273	- 142.612
r =		0.993	0.996	0.983

\* Data used for regression analysis.

Table - 10E

Effect of DD - 15 on growth of Helminthosporium oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	90.42*	84.92*
400	100	86.33*	82.14*	76.77*
300	100	75.27*	72.61*	63.25*
250	100	67.50*	63.50*	56.34*
200	100	59.05*	55.90*	47.50*
150	85.00	43.81*	42.15*	35.44*
100	71.15	31.25*	26.50*	20.02*
50	49.90	18.15	11.50	8.25
ED <sub>95</sub> =	> 150.00	500.00	524.80	616.59
ED <sub>50</sub> =	> 50.00	154.88	173.73	208.92
Regression constants: $Y = mx + c$				
m =		91.222	93.136	94.469
c =		-150.415	-159.243	-169.407
r =		0.999	0.998	0.999
* Data used for regression analysis.				

Table - 10F  
Effect of SD - 16 on growth of *Helminthosporium oryzae*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	89.09*	86.95*	78.51*
400	100	87.27*	81.52*	73.55*
300	100	78.50*	71.73*	65.23*
250	100	70.90*	63.04*	57.02*
200	100	64.50*	56.52*	48.76*
150	100	58.90*	48.50*	36.36*
100	76.20	45.45*	34.78*	27.27*
50	52.15	25.45*	21.25	15.15
ED <sub>95</sub> =	> 100.00	549.54	602.55	758.57
ED <sub>50</sub> =	< 50.00	114.81	158.43	199.52
Regression constants: $Y = mx + c$				
m =	0.00730	65.628	76.523	78.519
c =		-85.198	-118.408	-131.178
r =		0.997	0.998	0.995

\* Data used for regression analysis.

Table - 106

Effect of BD - 17 on growth of Helminthosporium oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	87.50*	77.96*
400	100	83.33*	78.75*	68.50*
300	100	68.50*	57.50*	55.93*
250	100	62.50*	50.00*	47.45*
200	100	52.75*	40.00*	32.93*
150	70.53*	38.50*	25.00*	23.72*
100	62.70*	23.50*	17.50*	14.10
50	41.17*	16.15	10.00	6.50
ED <sub>95</sub> =	343.73	524.80	602.55	707.94
ED <sub>50</sub> =	66.06	186.20	229.08	257.03
Regression constants: $Y = mx + c$				
$m =$	62.269	98.681	106.689	102.948
$c =$	- 62.483	- 174.025	- 202.417	- 193.760
$r =$	0.995	0.999	0.986	0.999
* Data used for regression analysis.				

Table - 101

Effect of ED - 18 on growth of Helminthosporium oryzae

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	78.90*	68.50*	62.50*
400	100	68.74*	58.61*	53.75*
300	68.42*	58.06*	47.50*	41.09*
250	60.90*	49.75*	38.46*	32.61*
200	50.75*	39.15*	30.75*	23.75*
150	36.50*	26.26*	17.07*	10.80*
100	20.57*	10.80*	8.75	5.25
50	14.42	6.65	5.50	0.00

ED<sub>95</sub> = 524.80

767.94

891.25

1023.20

ED<sub>50</sub> = 194.08

251.18

316.22

383.07

Regression constants:  $Y = mx + c$

$m = 103.237$

99.177

98.593

99.680

$c = -186.508$

-189.096

-196.446

-205.471

$r = 0.999$

0.999

0.999

0.999

\* Data used for regression analysis.

Table - 10I

Effect of Ninosan on growth of Helminthosporium oryzae.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after			
	24 hours	48 hours	72 hours	96 hours
500	100	100	100	100
400	100	100	100	100
300	100	100	97.50*	91.17*
250	100	100	90.47*	86.76*
200	100	91.83*	83.33*	77.94*
150	88.50*	79.59*	73.80*	67.64*
100	75.00*	65.30*	59.52*	52.50*
50	56.00*	40.50*	29.50*	24.50*
ED <sub>95</sub> =	186.20	218.77	269.15	309.02
ED <sub>50</sub> =	39.81	63.09	81.28	93.32
Regression constants: $Y = mx + c$				
m =	66.924	83.583	86.126	86.554
c =	57.559	-101.205	114.540	-121.071
r =	0.997	0.999	0.998	0.999
* Data used for regression analysis.				

Table - 11A

Effect of ED - 10 on spore germination inhibition of  
Aspergillus niger.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % inhibition	48 hours % inhibition
2000	100	64*
1600	71*	55*
1200	53*	39*
1000	44*	31*
800	31*	9
600	17*	0
400	0	0
$ED_{95} =$	2454.70	3630.78
$ED_{50} =$	1096.47	1445.43
Regression constants : $Y = mx + c$		
$m =$	125.999	111.737
$c =$	-333.285	-303.884
$r =$	0.998	0.998
* Data used for regression analysis.		

Table - 11B

Effect of ED - 11 on spore germination of *Aspergillus niger*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	93*	74*
1600	70*	57*
1200	45*	35*
1000	26*	0
800	7	0
600	0	0
400	0	0
ED <sub>95</sub> =	1995.26	2630.26
ED <sub>50</sub> =	1258.92	1412.53
Regression constants : $Y = mx + c$		
m =	218.071	169.548
c =	-626.790	-485.527
r =	0.998	0.999

\* Data used for regression analysis.

Table - 11C

Effect of DD - 13 on spore germination inhibition of *Aspergillus niger*

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	90*	75*
1600	78*	64*
1200	66*	46*
1000	60*	37*
800	49*	26*
600	37*	0
400	23*	0
ED <sub>95</sub> =	2290.86	2818.38
ED <sub>50</sub> =	776.24	1230.26
Regression constants : $Y = mx + c$		
m =	95.502	126.178
c =	-226.752	-337.702
r =	0.998	0.999
* Data used for regression analysis.		

Table - 11B

Effect of BD - 14 on spore germination inhibition of *Aspergillus niger*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	87*	75*
1600	78*	63*
1200	66*	49*
1000	60*	40*
800	49*	31*
600	37*	0
400	23*	0
ED <sub>95</sub> =	2398.83	3019.95
ED <sub>50</sub> =	794.32	1174.89
Regression constants: $Y = mx + c$		
$m =$	92.767	110.980
$c =$	-219.038	-201.775
$r =$	0.999	0.998
* Data used for regression analysis.		

Table - 11E

Effect of ED - 15 on spore germination inhibition of Aspergillus niger.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	77*
1600	85*	61*
1200	69*	49*
1000	58*	37*
800	44*	23*
600	25*	0
400	0	0
ED <sub>95</sub> =	1819.70	2754.22
ED <sub>50</sub> =	870.96	1230.26
Regression constants : $Y = mx + c$		
m =	140.567	131.175
c =	-363.816	-356.45
r =	0.999	0.995

\* Data used for regression analysis.

Table - III

Effect of BD - 16 on spore germination inhibition of Aspergillus Niger

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	65*	52*
1600	50*	40*
1200	33*	21*
1000	21*	11*
800	0	.0
600	0	0
400	0	0

ED<sub>95</sub> = 3162.27

ED<sub>50</sub> = 1548.81

Regression constants :  $Y = mx + c$

m = 143.868

c = -410.170

r = 0.958

\* Data used for regression analysis.

3981.07

1905.46

137.866

-402.246

0.999

Table - 11G

Effect of BD - 17 on spore germination inhibition of *Aspergillus niger*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	100
1600	100	81*
1200	76*	62*
1000	62*	49*
800	47*	33*
600	21*	12*
400	0	0
ED <sub>95</sub>	1479.10	1305.46
ED <sub>50</sub>	831.76	1000.00
Regression constants : $Y = mx + c$		
m =	180.353	161.785
c =	-477.837	-436.015
r =	0.993	0.999
* Data used for regression analysis.		

Table - III

Effect of ED - 18 on spore germination inhibition of *Aspergillus niger*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
2000	73*	69*
1600	69*	58*
1200	54*	44*
1000	44*	34*
800	33*	26*
600	21*	0
400	0	0

ED<sub>95</sub> = 2754.22

ED<sub>50</sub> = 1036.47

Regression constants :  $Y = mx + c$

m = 110.818

c = -227.503

r = 0.998

\* Data used for regression analysis.

3383.44

1318.25

102.668

-203.734

0.997

Table - III

Effect of Hincosam on spore germination inhibition of *Aspergillus niger*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	75*	53*
1600	66*	42*
1200	53*	29*
1000	47*	22*
800	41*	0
600	28*	0
400	11*	0

ED<sub>95</sub> = 3311.31

ED<sub>50</sub> = 1047.12

5123.61

1862.03

Regression constants :  $Y = mx + c$

m = 89.814

102.934

c = -221.533

-286.971

r = 0.999

0.999

\* Data used for regression analysis.

Table - 12A

Effect of ED - 10 on spore germination inhibition of Penicillium gansenii.

Concentration ( g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	81*
1600	88*	60*
1200	61*	43*
1000	54*	31*
800	41*	18*
600	23*	0
400	0	0
ED <sub>95</sub> =	1862.08	2511.88
ED <sub>50</sub> =	912.01	1288.24
Regression constants : $Y = mx + c$		
m =	146.489	154.964
c =	- 384.252	-432.859
r =	0.992	0.976
* Data used for regression analysis.		

Table - 12B

Effect of BD - 11 on spore germination inhibition of *Penicillium gensei*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	68*	57*
1600	59*	39*
1200	33*	17*
1000	23*	4*
800	0	0
600	0	0
400	0	0
ED <sub>95</sub> =	2951.20	3235.93
ED <sub>50</sub> =	1513.56	1819.70
Regression constants : $Y = mx + c$		
m =	150.675	175.454
c =	- 429.247	-522.114
r =	0.999	0.999

\* Data used for regression analysis.

Table - 120

Effect of BD - 13 on spore germination inhibition of *Penicillium gansenii*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	100
1600	100	95*
1200	92*	71*
1000	79*	65*
800	66*	38*
600	45*	14*
400	18*	0
$ED_{95} =$	1230.26	1543.31
$ED_{50} =$	630.95	891.25
Regression constants ; $Y = mx + c$		
$m =$	155.601	190.007
$c =$	-386.284	-511.142
$r =$	0.999	0.993
* Data used for regression analysis.		

Table - 122

Effect of BD - 14 on spore germination inhibition of *Penicillium saourei*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	70*	53*
1600	55*	39*
1200	37*	24*
1000	23*	13*
800	9*	0
600	0	0
400	0	0
	0	0
ED <sub>95</sub> =	2824.03	4168.59
ED <sub>50</sub> =	1445.43	1862.03
Regression constants : $Y = mx + c$		
m =	153.435	130.367
c =	-435.923	-337.431
r =	0.998	0.998

\* Data used for regression analysis.

Table - 123

Effect of BD - 15 on spore germination inhibition of Penicillium roseopurpureum.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	91*
1600	100	76*
1200	100	52*
1000	66*	41*
800	44*	32*
600	29*	28*
400	11*	8
	0	0

ED<sub>95</sub> = 1778.27

ED<sub>50</sub> = 812.83

Regression constants :  $Y = ax + c$

a = 132.757

c = - 338.543

r = 0.984

\* Data used for regression analysis.

Table - 12FEffect of ED - 16 on spore germination inhibition of Penicillium gamseni.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	69*	51*
1600	55*	37*
1200	38*	15*
1000	31*	6*
800	17*	0
600	0	0
400	0	0
ED <sub>95</sub> =	3162.27	3801.89
ED <sub>50</sub> =	1412.53	1949.84
Regression constants : $Y = mx + c$		
m =	123.276	153.097
c =	-354.887	-453.853
r =	0.999	0.998

\* Data used for regression analysis.

Table - 126

Effect of BD - 17 on spore germination inhibition of *Penicillium gensei*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	100
1600	100	77*
1200	77*	57*
1000	64*	43*
800	52*	32*
600	34*	17*
400	0	0
BD <sub>95</sub> =	1584.89	2137.96
BD <sub>50</sub> =	758.57	1023.29
Regression constants : $Y = mx + c$		
m =	139.980	140.567
c =	-354.082	-373.816
r =	0.997	0.998

\* Data used for regression analysis.

Table - 12H

Effect of BD - 18 on spore germination inhibition of *Penicillium gansenii*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	70*
1600	71*	66*
1200	53*	34*
1000	42*	24*
800	32*	0
600	11*	0
400	0	0
ED <sub>95</sub> =	2344.22	2630.26
ED <sub>50</sub> =	1096.47	1412.53
Regression constants : $Y = mx + c$		
m =	137.168	160.464
c =	-368.059	-474.615
r =	0.998	0.978
* Data used for regression analysis.		

Table 121Effect of Ninosan on spore germination inhibition of *Penicillium gensei*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	100
1600	100	71*
1200	68*	47*
1000	53*	32*
800	39*	18*
600	29*	8*
400	11*	0
ED <sub>95</sub> =	2187.76	2454.70
ED <sub>50</sub> =	891.25	1202.26
Regression constants : $Y = mx + c$		
m =	116.083	148.244
c =	-250.060	-407.553
r =	0.986	0.983
* Data used for regression analysis		

Table - 13AEffect of BD - 10 on spore germination inhibition of *Verticillium albo-atrum*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	57*	36*
1600	44*	24*
1200	31*	11*
1000	24*	0
800	13*	0
600	0	0
400	0	0
BD <sub>95</sub> =	4570.88	6918.30
BD <sub>50</sub> =	1737.80	2691.58
Regression constants : $Y = mx + c$		
m =	108.061	108.270
c =	-300.643	-321.716
r =	0.998	0.998
* Data used for regression analysis.		

Table - 13B

Effect of BD - 11 on spore germination inhibition of Verticillium albo-atrum.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	73*
1600	69*	55*
1200	43*	29*
1000	31*	17*
800	18*	0
600	0	0
400	0	0
ED <sub>95</sub> =	2290.86	2393.83
ED <sub>50</sub> =	1230.26	1445.43
Regression constants : $Y = mx + c$		
m =	171.316	203.493
c =	-430.979	- 594.737
r =	0.994	0.993

\* Data used for regression analysis.

Table 128

Effect of BD - 13 on spore germination inhibition of Verticillium albo-atrum.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	100
1600	100	85*
1200	88*	62*
1000	79*	49*
800	66*	37*
600	43*	14*
400	15*	
BD <sub>95</sub> =	1253.32	1819.70
BD <sub>50</sub> =	630.96	977.23
Regression constants: $Y = mx + c$		
m =	151.251	162.345
c =	-374.658	-435.689
r =	0.990	0.998

\* Data used for regression analysis.

Table - 192

Effect of ED<sub>50</sub> - 14 on spore germination inhibition of *Verticillium albo-atrum*.

Concentration ( $\mu$ g/ml)	Percentage of inhibition over control after	
	24 hours	48 hours
2000	100	71*
1600	95*	58*
1200	75*	44*
1000	67*	36*
800	50*	22*
600	32*	8*
400	0	0

ED<sub>95</sub> = 1594.89 3162.27

ED<sub>50</sub> = 776.24 1315.25

Regression analysis :  $Y = mx + c$   
 $m = 146.737$  115.916  
 $c = -374.800$  -321.671  
 $r = 0.090$  0.099

\* Data used for regression analysis.

Table - 13E

Effect of ED - 15 on spore germination inhibition of Verticillium albo-atrum.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	62*
1000	63*	47*
1200	48*	29*
1000	37*	20*
800	22*	8*
600	14*	0
400	0	0
ED <sub>95</sub> <sup>=</sup>	3019.95	3548.13
ED <sub>50</sub> <sup>=</sup>	1230.76	1621.81
Regression constants : $\bar{Y} = mx + c$		
m =	114.130	135.285
c =	-302.991	-385.374
r =	0.997	0.999

\* Data used for regression analysis.

Table - 13F

Effect of BD - 16 on spore germination inhibition of *Vorticillium albo-atrum*.

Concentration ( $\mu$ g/mL)	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	68*
1600	57*	39*
1200	41*	17*
1000	36*	0
800	22*	0
600	0	0
400	0	0
ED <sub>95</sub> =	3388.44	3235.93
ED <sub>50</sub> =	1342.96	1778.27
Regression constants : $Y = mx + c$		
m =	114.105	177.819
c =	-308.167	-529.244
r =	0.995	0.999
* Data used for regression analysis.		

Table - 130

Effect of BD - 17 on ~~growth~~ spore germination inhibition of Verticillium albo-atrum

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	88*	77*
1600	72*	60*
1200	57*	41*
1000	47*	29*
800	34*	14*
600	19*	0
400	0	0
ED <sub>95</sub>	2230.86	2570.39
ED <sub>50</sub>	1023.29	1348.96
Regression constants : $Y = mx + c$		
m =	129.330	156.830
c =	-340.331	-441.034
r =	0.998	0.999
* Data used for regression analysis.		

Table 13H

Effect of ED<sub>50</sub> 13 on spore germination inhibition of *Verticillium albo-atrum*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	60*	50*
1600	45*	35*
1200	27*	19*
1000	19*	0
800	7*	0
600	0	0
400	0	0
ED <sub>95</sub> =	3715.35	4265.74
ED <sub>50</sub> =	1698.24	1995.26
Regression constants : $Y = mx + c$		
m =	132.426	134.210
c =	-378.127	-393.464
r =	0.998	0.998
* Data used for regression analysis.		

Table - 181

Effect of Hinosen on spore germination inhibition of *Verticillium albo-atrum*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	100
1600	100	63*
1200	86*	52*
1000	87*	44*
800	72*	37*
600	59*	26*
400	43*	0
ED <sub>95</sub>	1174.89	3715.35
ED <sub>50</sub>	487.73	1122.01
Regression constants : $Y = mx + c$		
m =	112.837	86.141
c =	-252.218	-212.989
r =	0.995	0.998
* Data used for regression analysis.		

Table - 14AEffect of BD - 10 on spore germination inhibition of *Helminthosporium oryzae*.

Concentration ( $\mu\text{g}/\text{mL}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	100
1600	98*	89*
1200	75*	63*
1000	66*	49*
800	45*	28*
600	26*	7
400	0	0
ED <sub>95</sub>	1513.56	1659.58
ED <sub>50</sub>	812.83	1023.29
Regression constants : $Y = mx + c$		
m =	168.939	200.966
c =	-442.790	-579.031
r =	0.993	0.999
* Data used for regression analysis.		

Table - 14B

Effect of ED - 11 on spore germination inhibition of *Helminthosporium oryzae*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	98*
1600	88*	76*
1200	59*	40*
1000	44*	26*
800	26*	0
600	11	0
400	0	0
ED <sub>95</sub>	1698.24	1905.46
ED <sub>50</sub>	1047.12	1258.92
Regression constants : $Y = mx + c$		
m =	207.813	245.924
c =	-578.022	-712.817
r =	0.998	0.998

\* Data used for regression analysis.

Effect of BD - 13 on spore germination inhibition of Helminthosporium oryzae.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	84*
1600	75*	70*
1200	56*	41*
1000	42*	28*
800	27*	10*
600	11*	0
400	0	0
$\text{ED}_{95}$	2137.96	2187.76
$\text{ED}_{50}$	1071.51	1288.24
Regression constants : $Y = mx + c$		
m =	151.494	190.309
c =	-410.464	-542.218
r =	0.997	0.997

\* Data used for regression analysis.

Table - 14B

Effect of BD - 14 on spore germination inhibition of Helminthosporium oryzae.

Concentration ( $\mu$ g/ml.)	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	100
1600	100	100
1200	92*	81*
1000	73*	63*
800	51*	45*
600	24*	16*
400	0	0
ED <sub>95</sub>	1230.26	1380.38
ED <sub>50</sub>	776.94	831.76
Regression constants : $Y = mx + c$		
m =	224.361	212.475
c =	-593.500	-572.365
r =	0.998	0.998
* Data used for regression analysis.		

Table - 14EEffect of ED - 15 on spore germination inhibition of *Helminthosporium oryzae*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	76*
1600	72*	60*
1200	56*	43*
1000	44*	29*
800	32*	16*
600	17*	0
400	0	0
ED <sub>95</sub>	2393.83	2630.26
ED <sub>50</sub>	1071.51	1318.25
Regression constants : $Y = mx + c$		
m =	129.464	150.456
c =	-342.610	-420.713
r =	0.998	0.998

\* Data used for regression analysis.

Table - 14<sup>a</sup>Effect of BD - 16 on spore germination inhibition of Helminthosporium oryzae

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	76*
1600	69*	53*
1200	53*	34*
1000	40*	24*
800	31*	0
600	11*	0
400	0	0
ED <sub>95</sub>	2454.71	2570.39
ED <sub>50</sub>	1122.01	1412.53
Regression constants : $Y = mx + c$		
m =	133.806	175.500
c =	-359.013	-503.510
r =	0.936	0.939
* Data used for regression analysis.		

Table - 14G

Effect of BD-17 on spore germination inhibition of *Helminthosporium oryzae*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	100
1600	100	100
1200	87*	63*
1000	73*	48*
800	56*	36*
600	25*	19*
400	0	0
ED <sub>95</sub>	1258.92	2041.73
ED <sub>50</sub>	758.57	977.23
Regression constants : $Y = mx + c$		
m =	204.420	142.043
c =	-539.723	-375.396
r =	0.997	0.993
* Data used for regression analysis.		

Effect of BD - 18 on spore germination inhibition of *Helminthosporium oryzae*.

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours	48 hours
	% Inhibition	% Inhibition
2000	100	100
1600	100	83*
1200	65*	57*
1000	49*	45*
800	29*	20*
600	8*	0
400	0	0

BD <sub>95</sub>	1693.24	1778.27
BD <sub>50</sub>	977.23	1071.51

Regression constants :  $Y = mx + c$

m =	189.096	207.194
c =	-517.247	-579.107
r =	0.997	0.997

\* Data used for regression analysis.

Table - 141

Effect of Kinogen on spore germination inhibition of *Helminthosporium oryzae*

Concentration ( $\mu\text{g/ml}$ )	Percentage of inhibition over control after	
	24 hours % Inhibition	48 hours % Inhibition
2000	100	100
1600	100	100
1200	100	100
1000	97*	86*
800	88*	69*
600	72*	49*
400	57*	23*
$ED_{95}$	933.25	1148.15
$ED_{50}$	338.89	588.84
Regression constants : $Y = mx + c$		
$m =$	101.979	156.760
$c =$	-208.826	-384.922
$r =$	0.977	0.999
* Data used for regression analysis.		

