



Root rots caused by Phycomycetes

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BY PHYCOMYCETES

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CHRISTINE J. BUISMAN

ROOT ROTS CAUSED BY
PHYCOMYCETES

CHRISTINE JOHANN
BLUMEN

UNIVERSITEITSBIBLIOTHEEK UTRECHT



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ROOT ROTS CAUSED BY PHYCOMYCETES

PROEFSCHRIFT

TER VERKRIJGING VAN DEN GRAAD VAN
DOCTOR IN DE WIS- EN NATUURKUNDE
AAN DE RIJKS-UNIVERSITEIT TE UTRECHT,
OP GEZAG VAN DEN RECTOR MAGNIFICUS
Dr. A. NOORDT'ZIJ, HOOGLEERAAR IN DE
FACULTEIT DER GODGELEERDHEID,
VOLGENS BESLUIT VAN DEN SENAAT DER
UNIVERSITEIT TEGEN DE BEDENKINGEN
VAN DE FACULTEIT DER WIS- EN NATUUR-
KUNDE TE VERDEDIGEN, OP DINSDAG
22 MAART 1927, DES NAMIDDAGS VIER UUR

DOOR

CHRISTINE JOHANNA
BUISMAN

GEBOREN TE LEEUWARDEN

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INTRODUCTION

These investigations were originally undertaken with the idea of adding something to our knowledge of the root rot of peas caused by *Phycomycetes*. In America several publications have appeared about it during the last few years, while very little attention has up to now been given to it in Europe. It would, for instance, be of importance to investigate if the dangerous american root parasite of peas, *Aphanomyces euteiches*, Drechsler, is also present in diseased pea roots in this country and if it is the cause of rootrot.

As it was necessary that I first gained some experience with *Phycomycetes*, I examined several different diseased plants which were given to me as probably being infected by a fungus of that group.

I will also discuss some of these diseases which up till now were still unknown.

This applies in the first instance to the disease of the roots of the *Calla-lily* which causes a considerable loss to the *Calla-nurseries* in this country and has been thought to be due to *Nematodes*. It happened that diseased *Chrysanthemum* cuttings, which had suddenly stopped growing, were also sent to the laboratory at the time.

In the course of time I could examine different rots of cuttings, several of which I described in this publication.

While examining the different pea roots it struck me, that *Asterocystis radidis* de Wildeman, the wellknown parasite on flax roots which is taken to be the cause of „brand“ = fire in flax, is often found on them. In order to be able to recognize this *Chytridiacea* and not to mistake it for oogonia of *Phycomycetes* I also examined roots of diseased flax. In doing so I discovered a

Phycomycete which is apparently parasitic on the flax roots, and which I have therefore also described here.

As most of these plants had to be cultivated in vegetable mould it was necessary to know something about the *Phycomycetes* which inhabit humus. If our soil proved to be strongly infected by them, inoculation experiments in it would be worthless. Some more observations about the different *Phycomycetes* which I isolated from seedlings grown in our vegetable mould will also have to be considered.

This work was carried out more along phytopathological than along morphological lines. I could only make a choice out of all the isolations which were at my disposition.

A thorough treatment of the genera *Pythium* and *Phytophthora* would lead me too far, so I am restricting myself to some considerations about the classification of the group to which they belong, and about some related genera.

CHAPTER I

TAXONOMY

§ 1. *The classification in the Phytophthoreae*

In the year 1881 de Bary (4) already wrote about the classification of the group *Phytophthoreae*, in connection with the genera *Pythium* and *Phytophthora*. He showed that there is far too little difference between these genera to justify placing them in different families as Pringsheim had done (*Pythium* with the *Saprolegniaceae*, *Phytophthora* with the *Peronosporaceae*). De Bary pointed out, that there is really no other difference between *Pythium* and *Phytophthora* than the way in which the zoospores are formed. In the genus *Pythium* the protoplasm from the sporangium flows into a vesicle in which the zoospores are then formed, whereas in the genus *Phytophthora* the zoospores are formed in the sporangium itself.

Butler's monograph (10) about the genus *Pythium* is the best source of data at present. He agrees with de Bary and also sees no other difference between *Pythium* and *Phytophthora*. He includes both genera in the *Peronosporaceae*.

A detailed discussion about this question is found in a chapter from „Generic concepts in the *Pythiaceae* and *Blastocladiaceae*” by Fitzpatrick (14), i. e. *The Pythium — Phytophthora Problem*.

This shows, that an absolute separation between *Pythium* and *Phytophthora* is theoretically impossible, as it has become evident that the traditional characteristic which separates *Pythium* from *Phytophthora*, namely the formation of zoospores in a vesicle, in-

stead of in the sporangium, is also found amongst different *Phytophthora* species. —

But in literature we find other species which belong to the group of *Pythium* and *Phytophthora*. How are they related to *Pythium* and *Phytophthora*?

In the first place the genus *Pythiacystis*. Barrett (2), E. Smith and R. Smith (33) and Leonian (20) pointed out, that a separation between *Pythiacystis* and *Phytophthora* cannot be carried through. Smith and Smith even isolated forms, which showed a transition from *Pythiacystis* to *Phytophthora cactorum*. Leonian has now definitely dropped the genus *Pythiacystis* and uses the name *Phytophthora citrophthora*.

Dufrénoy (13) includes the genus *Blepharospora* Petri in the homogenous group of *Phytophthora*, *Pythium* and *Pythiacystis* (which he, like Fitzpatrick (14), separates from the real *Pero- nosporaceae* with aerial conidia and which are obligate parasites).

Fitzpatrick further regards the genus *Pythiogeton* v. Minden as a representative of this group.

According to my opinion, *Pythiomorpha* Petersen is another genus which belongs to it.

The question is, whether there are sufficient reasons for letting these genera exist next to *Pythium* and *Phytophthora*. I do not think, that there ought to be any difficulty in letting the genus *Pythiogeton* remain, as it is sufficiently characterized by its asymmetrical sporangia. — It is a different question with *Blepharospora* and *Pythiomorpha* however. —

According to my opinion these genera, each with one representative, *Blepharospora cambivora* Petri and *Pythiomorpha gonapodyides* Petersen, should be included in the genus *Phytophthora*. The reasons for this I will give below.

I do not think that Petri (27, 28, 29) succeeded in proving definitely that his *Blepharospora* is not a *Phytophthora*.

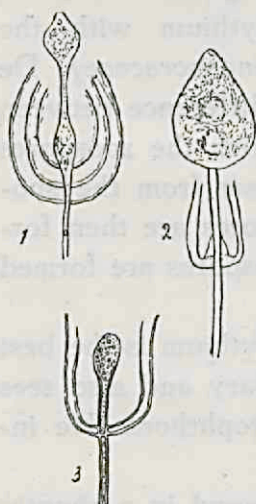


Fig. 1

1 and 2: Zoosporangium of *Pythiomorpha gonapodyides* after Miss Kanouse (high power).

3: Zoosporangium of *Blepharospora cambivora* after Petri. $\pm 333\times$

His principal arguments in favour of making a new genus are:

1. that *Blepharospora* does not produce sporangia on the surface of the infected parts of the plants or on solid culture media, but only in a mineral salt solution.
2. that *Blepharospora* possesses long, unbranched sporangio-phores ending in one sporangium.

It will become obvious below that representatives of the genus *Phytophthora* also show similar characteristics.

Blepharospora is related to *P. cactorum*, *P. fagi* and *P. syringae*, according to Petri, because it possesses paragynous antheridia. Although Pethybridge does no longer consider the possession of paragynous or amphigynous antheridia a reason for subdividing the genus *Phytophthora*, as both kinds of antheridia sometimes occur in one form, Petri still maintains that this subdivision should remain.

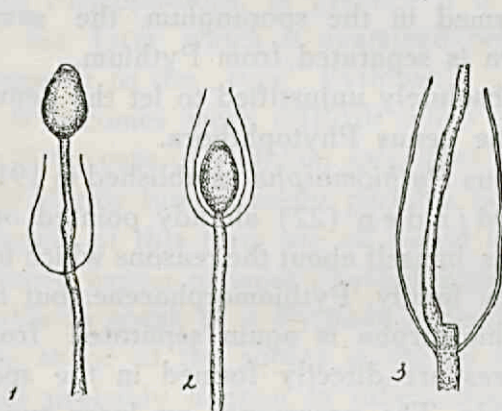


Fig. 2

Zoosporangia of *Phytophthora cinnamomi*
after Rands.

1 and 2 $\pm 237 \times$, 3 $370 \times$

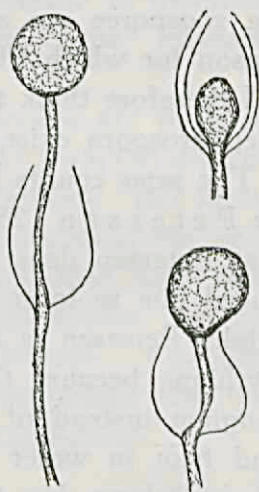


Fig. 3

Zoosporangia of *Phytophthora cryptogea* after
Pethybridge. $340 \times$

A *Phytophthora* would then belong to the one or to the other group, as it more regularly produces amphigynous or paragynous antheridia. In this case *Blepharospora* would then belong close to *Phytophthora cactorum*. I do not attach very much importance to this production of antheridia, as *Blepharospora* forms oogonia only in the tissues of the infected chestnut-tree, and it is therefore

difficult to determine the position of the antheridium. The characteristics show that *Blepharospora* is more closely related to *Phytophthora cryptogea* and *P. cinnamomi*.

These characteristics are exactly the same as the ones, Petri uses to separate *Blepharospora* from the genus *Phytophthora*, namely the difficulty in forming sporangia and the unbranched sporangiophores.

Phytophthora cryptogea and *P. cinnamomi* also rarely produce sporangia and possess little branched conidiophores. Furthermore these two genera also have another characteristic in which they correspond with *Blepharospora*. The sporangiophore grows into the sporangium after its contents have been discharged. The sporangia are also rounded off above, without a definite papilla being formed, exactly as is the case with *Blepharospora*.

Petri separates *Blepharospora* from the genus *Pythium* because the zoospores are already formed in the sporangium, the same reason for which *Phytophthora* is separated from *Pythium*.

I therefore think that it is absolutely unjustified to let the genus *Blepharospora* exist next to the genus *Phytophthora*.

The same counts for the genus *Pythiomorpha* established in 1910 by Petersen (25). Von Minden (22) already pointed out that Petersen does not express himself about the reasons which induced him to form even a new family, *Pythiomorphaceae*, but he thinks Petersen is right. *Pythiomorpha* is again separated from *Pythium*, because the zoospores are directly formed in the sporangium instead of in a vesicle. The occurrence on loose twigs and fruit in water is not a special characteristic of the genus *Phytophthora*, but this is no reason for excluding *Pythiomorpha* from the genus *Phytophthora*, as it is known (see Dufrénoy) (13) that *Phytophthora*'s may lead a saprophytic life.

In the detailed description of *Pythiomorpha gonapodyides*, the only existing species of the genus *Pythiomorpha*, by Miss Kanouse (19) there is no characteristic to be found which should separate *Pythiomorpha* from the *Phytophthora*'s. Petersen emphasized the fact that the mycelium is coloured purple by chlorojodide of zinc, but when I applied the same method to *Blepharospora*, it gave the same results.

Pythiomorpha also shows the sporangiophores which grow into

the sporangium after the swarmspores have been set free and sporangia and oogonia are formed with difficulty just as in *Blepharospora*.

I think that I have now shown that the genera *Blepharospora* and *Pythiomorpha* should be included in the genus *Phytophthora*, so that we may speak of *Phytophthora cambivora* and *Phytophthora gonapodyides*. Then except *Pythiogeton*, only the genera *Pythium* and *Phytophthora* remain, as *Pythiacystis* has already been included in the genus *Phytophthora*.

According to my opinion it would be of no practical importance to unite these two genera into one. The name *Phytophthora* is so universally used that it would be confusing to drop it in favour of the (older) name *Pythium*.

It is enough to state, as other investigators have already done, that theoretically no sharp line can be drawn between *Pythium* and *Phytophthora*. In practice it is usually possible to determine if the form which is examined belongs to the „type“ *Phytophthora* or to the „type“ *Pythium*.

It becomes more difficult when no swarmspores are produced as in the case of *Pythium splendens* Braun. Braun (8) does not say if this fungus might perhaps be a *Phytophthora*. But as the conidia of this form are all round instead of possessing the pear-shaped aspect of most *Phytophthora*-sporangia, it was only reasonable to speak of a *Pythium* instead of a *Phytophthora*, although the shape of the spores is not an official difference. The same will probably happen in other cases where difficulties are met with in connection with the production of swarmspores.

§ 2. The classification in the genera *Phytophthora* and *Pythium*

Few genera have been studied so often as *Phytophthora*, and yet the classification is far from being settled.

At present it is very difficult and sometimes nearly impossible, to identify a *Phytophthora*. The different species of this genus are so much alike, that it is nearly impossible to determine which species one is dealing with. From the old species of de Bary, *P. omnivora*, the four species *P. fagi*, *P. cactorum*, *P. syringae* and *P. faberi* have since been separated. Leonian (20) now states

that there is no difference between *P. fagi* and *P. cactorum*, and eliminates *P. fagi*.

There are many other difficulties in the same way. It was first tried to settle these questions along purely morphological lines, afterwards physiological methods were tried for the same purpose.

Rosenbaum (32) wrote a very accurate study in which he gives a detailed morphological account of the different species. He finally separates them by means of several characteristics, for instance the difference in size of the zoosporangia and oogonia as is indicated by curves.

But this is a very difficult method to apply in practice. This publication of Rosenbaum is however still a very valuable acquisition to the *Phytophthora* literature, especially as the species are studied on different culture-media. Leonian's research was meant to give a solution of the problem along physiological lines. He declares, specially in connection with his investigations about the mutations of *Phytophthora parasitica rhei*, that the genus may not be classified according to the size of the sporangia, the presence or absence of oogonia, other characteristics which appear in culture, and pathogenicity.

In practice there are however many objections against his „physiological key". Leonian himself also realizes that it is very difficult to cultivate fungi under exactly the same physiological circumstances in different parts of the world.

It therefore becomes nearly impossible to split the genus up into several divisions and to draw a sharp line round each. The matter has become more complicated by the publications of Ashby (1) and Gadd (15).

They demonstrated, that heterothallism is found in the species *Phytophthora faberi*. This could also show, that different species of *Phytophthora* may form hybrids in this way, and as transitions between the different forms will then of course originate, a sharp separation between them would become absolutely impossible.

According to my opinion, the morphological differences between the species must for the present first be determined and then physiological characteristics may be added, so that we may get a system which will be useful in practice. The more so, as it is

easier to identify by means of morphological characteristics, than by the round about physiological method.

In the genus *Pythium* things are not so complicated as with *Phytophthora*. Still it is not at all easy to determine if a special isolation belongs to the otherwise wellknown *Pythium de Baryanum* or not.

Even Drechsler (12) who is writing a monograph about the genus *Pythium* speaks of „species” or „forms” of the *de Baryanum* type. I think that we must now be specially careful not to give too strict definitions of the species, as the *Phycomycetes* are very variable forms. Some species of *Pythium* for instance show a different tendency towards forming oogonia or sporangia. De Bary (4) already noticed this in 1881 with *P. megalacanthum*. I possess only one isolation of this species which forms only oogonia and no sporangia.

The same is the case with an undescribed *Pythium* I isolated from various plants. The one isolation forms sporangia and swarm-spores, although only few, the other nearly purely oogonia.

Nothing further is known about the sexuality of these forms. — Perhaps the many forms one meets without oogonia show that heterothallism exists as in *P. faberi*, or perhaps we have here a strong tendency towards production of asexual spores. This appears to be the case with *P. de Baryanum* var. *geranii* Braun (8) which produced a few oogonia in culture only after 14 months which even then often degenerated.

I also came across this form when isolating, but never noticed any oogonia.

Some uncertainty therefore also exists where the genus *Pythium* is concerned.

I never noticed anything of biological races adapted to certain hosts. According to Drechsler however, some forms of *Pythium de Baryanum* are more pathogenic than other forms of the same species.

This again shows the great variability which exists, also in connection with physiological characteristics. It is therefore advisable not to give too strict a definition of the different species.

CHAPTER II

EXPERIMENTAL METHODS

In making isolations from infected roots of the different hosts I proceeded as follows:

The diseased roots were layed out on agarplates without being disinfected, as this handling would have killed the delicate mycelium of the *Phycomycetes*. At first I always used cherry-agar for the isolations as it is rather acid and therefore arrests growth of bacteria. Afterwards I found that oatmeal-agar is a more desirable medium. Although it is more favourable for bacterial growth, the *Phycomycetes* grow better on it than on cherry-agar.

It is very difficult to keep the cultures absolutely free from bacteria as they adhere to the hyphae and are therefore difficult to remove. The best way is to cultivate the fungi on agarplates and to transfer them as often as possible.

I usually used oatmeal-agar for keeping them in culture. On cherry-agar they mostly also produce a woolly growth of mycelium but few oogonia are formed. *Pythium de Baryanum* however grows well on cherry-agar while *P. megalacanthum* does not grow at all.

On cornmeal-agar many oogonia are also formed, but this medium is even more favourable for the growth of bacteria and therefore not so suitable.

Swarmsporangia are, on a whole, seldom produced on agar media. In order to obtain them I adopted the method used by Petri for *Blepharospira cambivora*.

I mostly used the receipt as given by him:

0,400 gr. $\text{Ca} (\text{NO}_3)_2$
 0,150 gr. KH_2PO_4
 0,150 gr. Mg SO_4
 0,060 gr. KCl .
 1000 gr. distilled water.

I cultivated a tuft of the mycelium from agarcultures for a few days in a flask, in which a thin layer of this solution had been brought. Hardly any growth of mycelium took place but apparently this anorganic solution served as a stimulus for the production of sporangia. The greatest number of sporangia were however formed when the bit of mycelium was afterwards transferred from the solution to a watchglass with distilled water. Sometimes the zoospores appeared directly; in other cases the water had to be refreshed first, and zoospores only appeared afterwards, for instance after one day. This method was successful with several kinds of *Phytophthora*-and *Pythium*-species which had at first hardly produced any zoospores, or none at all.

Even species which had been isolated years ago and had since then been kept in culture at the „Centraal Bureau voor Schimmelcultures“, could be induced to form zoospores when treated in this way. With *Aphanomyces euteiches* I also obtained the same results.

It is a pity that the results are not always so satisfactory: *Pythium splendens* for instance, cannot be induced to form zoospores even by this method. In other cases there are so few that it is impossible to trace their development.

For inoculation material I used some strains which I had isolated from diseased plants, and some species which were kept in culture at the „Centraal Bureau v. Schimmelcultures“. I never found any indications that the pathogenicity of these species had suffered by remaining in culture for so long. *Pythium complectens* from Braun, *Pythium splendens* from Leonian, and *P. de Baryanum* from Hartley which had been cultivated for some time, were still very virulent.

I will enter more fully into the methods of infection when I describe the experiments. The main point is that I often used watercultures in which the fungus was brought (see chapters on Calla-lily, peas and flax). As for the cuttings, their lower ends

were covered with the inoculation-material and then placed in sterilized sand.

Whenever I used vegetable mould for growing peas in, I sterilized it before use, because of the many *Phycomycetes* occurring in humus. (See chapter on Occurrence of *Pythium* in vegetable mould).

CHAPTER III

ROOT ROT OF THE CALLA-LILY

§ 1. *Occurrence of the disease*

Large hothouses with Calla-lilies are found on several commercial nurseries in our country. It is a very profitable culture as the flowers are very valuable and as each hothouse may yield some thousands of flowers.

The plants are treated in the following manner:

In spring they are put in the open, so that they may dry out and pass through a period of rest. After this period the corms are taken out of the pots, cleaned and planted again in fresh pots. They are left outside for another little while (not dry this time) and put in the hothouses at the beginning of winter. They start flowering in about December. In the last few years it often happened at several nurseries that the plants started to fade in December. The outer leaves then showed a yellow discoloration, which slowly spread over the surface until the whole leaf had turned yellow. The same happens to the other leaves until there are only a few left. The flowers are few and small so that the yield of a hothouse with diseased plants is very small. The disease is not very acute as new leaves are still being formed and the plant is not killed, but it remains ailing.

A diseased plant is easily withdrawn from the pots as the roots are easily broken off, and only small rotten pieces still adhere to the corm. The corm itself is sometimes also full of holes and grooves. If the plant is very carefully taken out of the pot with soil and

all, the roots appear to be badly infected. They have a glassy and drooping appearance, especially striking with the thicker roots. A reddish discoloration is also often found.

If the plants are treated in the usual way, e. g. if they pass through a period of rest and are then changed to other pots, there is at first no evidence of the disease. But, about the flowering season the symptoms appear again. It stands to reason that the corms suffer badly from such repeated occurrence of the disease.

§ 2. *Microscopical examination of the roots and isolation of the pathogenic organism*

One receives the same confused impression when examining the infected roots of the Calla-lily as when examining other diseased roots. First one notices the many *Nematodes*. The disease had at first been ascribed to them, but as most of them belonged to the not-parasitical kind *Rhabditis brevispina*, which lives in all kinds of decaying portions of plants, their presence was still no proof that they are indeed the cause of the disease.

Besides these *Nematodes* different kinds of soil-fungi are found in the roots. *Thielavia basicola* was occasionally found and also often oogonia of *Phycomycetes*. It was tried to find the cause by laying out small pieces of infected roots on agarplates. As the roots were so delicate, it was of course impossible to sterilize them first. This proved to be a great objection as all kinds of fast growing soil-fungi and saprophytes, which immediately attack the decaying roots, now appeared in my cultures. The isolations therefore only yielded common fungi such as *Pythium*-species and *Botrytis*. They are just ordinary inhabitants of vegetable mould and it was improbable that they would be the cause of the Calla-lily disease, which appeared relatively suddenly. After several unsuccessful attempts I gave up this method of isolating the pathogenic organism.

In the nurseries it had become evident that the disease also appears when the corms were dried and planted in absolutely fresh soil. This would show that the disease is transmitted by the corms.

It was therefore necessary to cultivate the corms under such circumstances that ordinary soil-fungi would be excluded. If the

corms then still became infected, it ought to be easier to isolate the pathogenic organism. As Calla-lilies under ordinary circumstances grow in marshes and swamps, it was probable that it would be possible to cultivate them in culture-solutions. As the big corms are awkward to work with I used very young ones in my experiments.

I took some young corms of infected plants, brushed half the number with water and the other half with a fairly strong solution of formalin (commercial solution diluted about 20 times). I then took a glass vesicle which had first been painted black and then white, so that the temperature of the liquid inside would not become too high. A lid, through which a few holes had been made, could be screwed on it. The vesicle was filled with culture solution of v. d. Crone:

1 L. water

1 gr. KNO_3

0,5 gr. Ca SO_4

0,5 gr. Mg SO_4

0,25 gr. $\text{Fe}_3 (\text{PO}_4)_2$

0,25 gr. $\text{Ca}_3 (\text{PO}_4)_2$.

The young corms were allowed to hang in the culture solution and were fastened in the holes in the lid by means of two pieces of cork, which fitted in each hole. In this way the leaves could freely develop into the air while the corms touched the culture-solution. After some time it became evident that the plants which had been treated with formalin, had developed excellently. The roots were strong and healthy. The roots of the plants which had only been treated with water, on the other hand, soon showed signs of infection. This rot could not be the result of an organism present in the solution, as the formalin-plants remained healthy.

When the infected roots of the plants in the solution were examined microscopically, they were found to be covered by a large number of sporangia, which soon formed swarmspores in water. Hardly any other fungi were present. When I layed out small pieces of these infected roots on agarplates the fungus only grew very slowly, although such a number of swarmsporangia were present.

I nevertheless succeeded in isolating it in this way as the usual soil-fungi were absent.

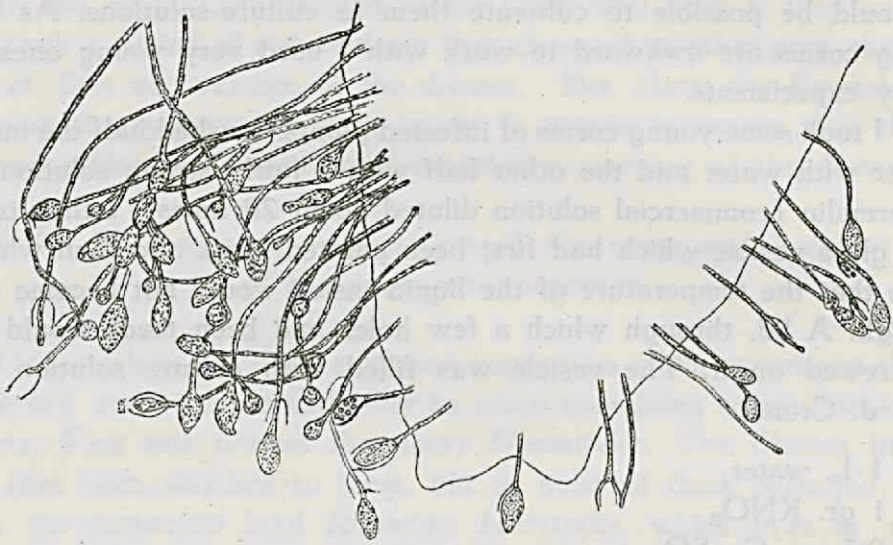


Fig. 4

Part of diseased Calla-root, crushed under the cover-slip.

$\pm 80 \times$

§ 3. *The pathogenic organism*

Inoculation experiments, which will be described below, proved, that the fungus with swarmsporangia which was found on the diseased roots was indeed the cause of the rot. I afterwards also found it on a root of a diseased plant which had not been grown in waterculture but had been taken out of the soil.

The way in which the swarmspores are formed could directly be followed when infected roots were examined in water. It appeared that this happens inside the sporangia in the way characteristic for *Phytophthora*.

It also became evident that, after the swarmspores have been set free, the sporangiophore grows into the sporangium in the same way as is found in the group of *Phytophthora cryptogea*, as I have already pointed out.

Little branched sporangiophores and sporangia without a distinct germinal papilla are also found.

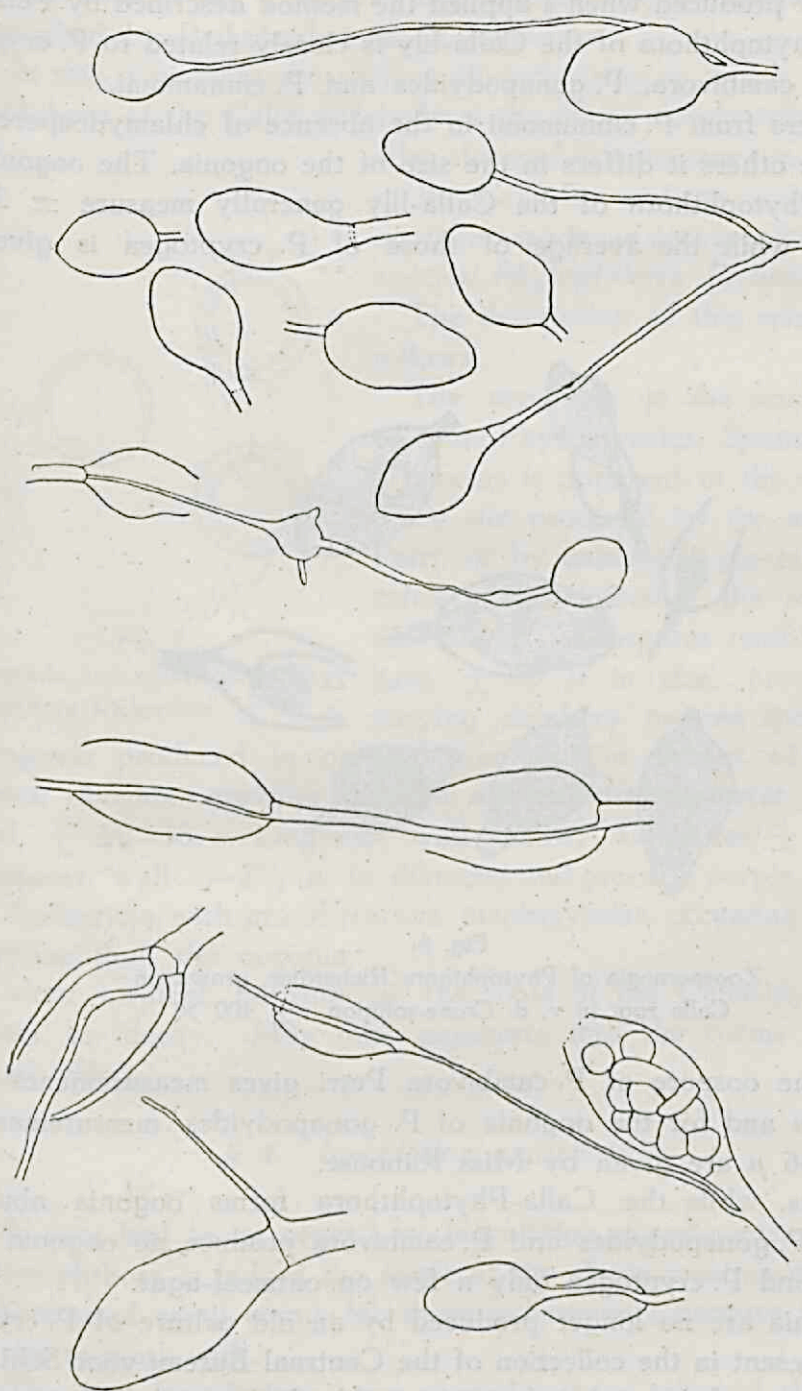


Fig. 5

Fig. 6
 Zoosporangia of *Phytophthora richardiae* in culture-solution.
 $\pm 366 \times$

Many oogonia with amphigynous antheridia were present in cultures on oatmeal-agar, but sporangia never appeared. These were only produced when I applied the method described by Petri.

The *Phytophthora* of the Calla-lily is closely related to *P. cryptogea*, *P. cambivora*, *P. gonapodyides* and *P. cinnamomi*.

It differs from *P. cinnamomi* in the absence of chlamydospores. From the others it differs in the size of the oogonia. The oogonia of the *Phytophthora* of the Calla-lily generally measure $\pm 34 - 38 \mu$, while the average of those of *P. cryptogea* is given as 30μ .

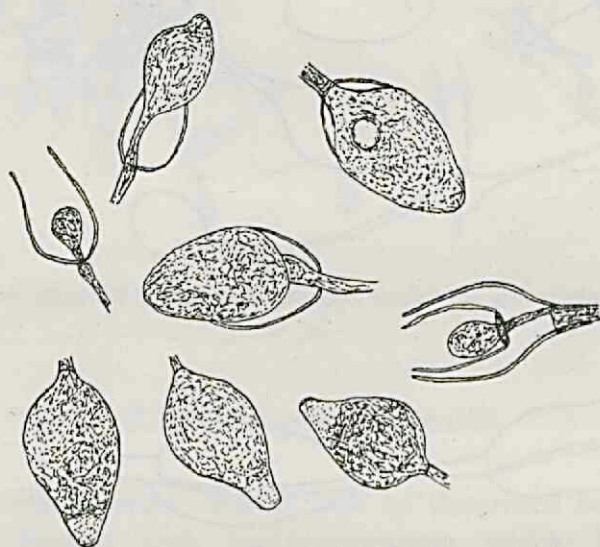


Fig. 6

Zoosporangia of *Phytophthora Richardiae*, grown on Calla root in v. d. Crone-solution. $\pm 400 \times$

For the oospore of *P. cambivora* Petri gives measurements of $20-27 \mu$ and for the oogonia of *P. gonapodyides* measurements of $22-36 \mu$ are given by Miss Kanouse.

Besides, while the Calla-*Phytophthora* forms oogonia abundantly, *P. gonapodyides* and *P. cambivora* produce no oogonia in culture and *P. cryptogea* only a few on oatmeal-agar.

Oogonia are no longer produced by an old culture of *P. cryptogea* present in the collection of the Centraal Bureau voor Schimmelcultures. It was therefore impossible for me to compare the

oogonia of the Calla-Phytophthora with those of the three last mentioned Phytophthora's. Nevertheless I think, that the colour of the wall of the oogonium and oospore also differs from those described by Pethybridge for *P. cryptogea*.

In this way there are various characteristics in which the Phytophthora of the Calla differs from the forms closely related to it.

Also the special occurrence in the roots of the Calla-lily and the virulent action on them, made me decide to form a new species *Phytophthora Richardiae*.

The description of this species is as follows:

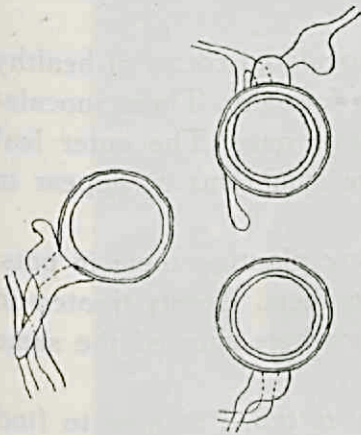


Fig. 7

Oogonia and antheridia of *Phytophthora Richardiae*. $\pm 366\times$

Oogonia produced in oatmeal-agar and in tissues of infected roots, obovate, growing through antheridium, diameter of distal end $\pm 34-38\ \mu$. Oogonial wall yellow. Oospores $\pm 29\ \mu$ in diameter, wall $2-2\frac{1}{2}\ \mu$. in diameter and greenish purple in colour.

Antheridia with protuberances, amphigynous, occurring on other hyphae than the oogonia.

Very virulent parasite of the roots of the Calla-lily causing them to decay. May also penetrate into the corms with the same effect.

§ 4. Inoculation experiments

It now had to be proved by inoculation experiments, that the *Phytophthora* is indeed the cause of the above mentioned disease. I first used small corms for these experiments because they are easier to work with.

The first inoculations were carried out on some of the young

corms treated with formalin. They were quite healthy and grew vigorously in the culture solution, although they had been taken from diseased plants. Small pieces of agar with *Phytophthora* mycelium were smeared on the roots. After a few days the roots began to show signs of decay, they became flabby and took on a glassy appearance.

This spread further and further until after some time all the roots were badly infected.

In order to be absolutely sure, I then inoculated corms of healthy plants which had also been brushed with formalin. These inoculations gave the same results as the previous ones. The outer leaf of an infected corm soon shows the same symptoms as appear in fullgrown diseased plants.

Finally I also infected large corms before planting them in pots, by rubbing a culture of *Phytophthora* on them. Corms treated in this way formed very few roots and on a whole showed the same symptoms of disease.

As the corms are sometimes only partly rotten, I wanted to find out if this was also due to infection by *Phytophthora*. I therefore laid slices of the corms in petridishes on wet filterpaper and inoculated them with the *Phytophthora*. A very peculiar rot, differing from the slimy bacterial rot, appeared. The cells seemed to become detached from each other and a fairly dry mass was formed.

§ 5. Control

It was now necessary to find a method of controlling the disease. As I had been successful in treating the corms with formalin it was probable that a solution to the problem could be found in this direction. Originally I had immersed the corms before planting them in a 2 0/0 solution of formalin and then brushed them with it.

As it appeared that this treatment was not sufficient, I think it better to leave them in a weaker solution for some time. The rotten places should be cut out.

Investigations of Bewley (5,6) already showed that the corms do not suffer by this treatment with formalin. Bewley reported an epidemic in Calla-lily nurseries, which seems to be very

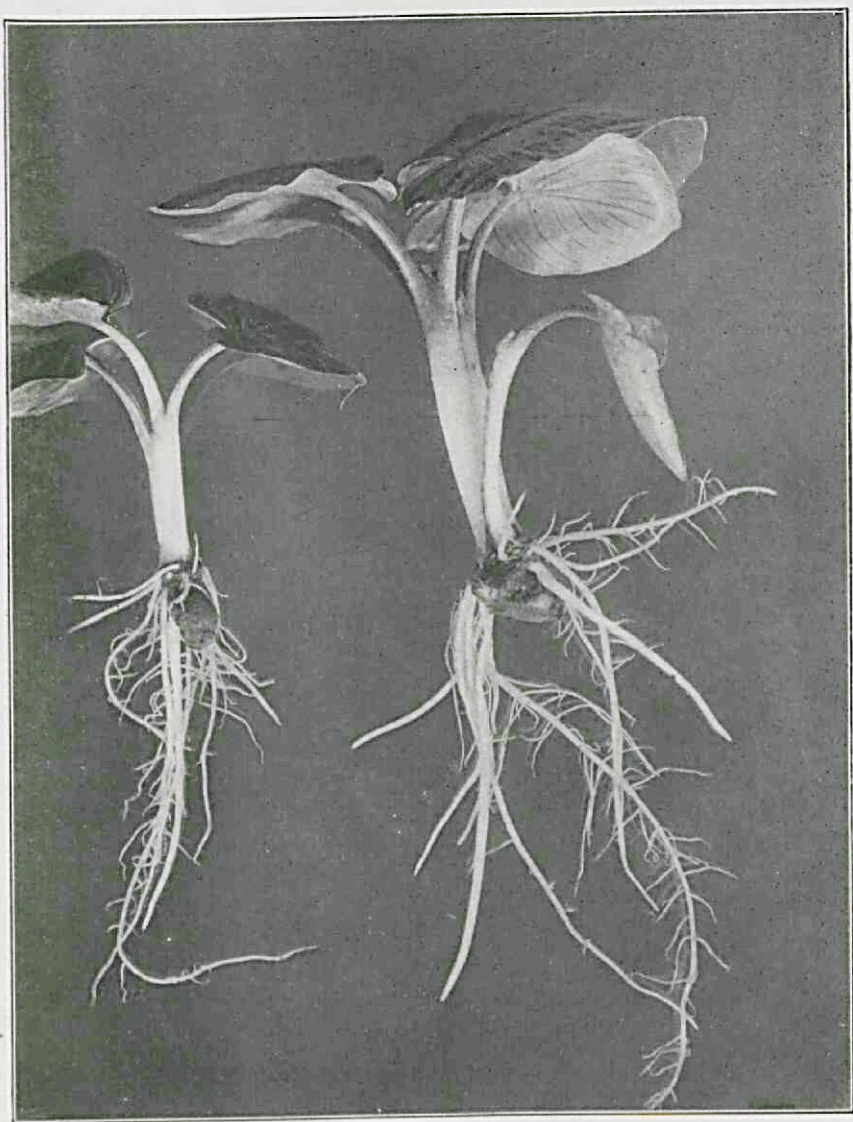


Plate I

Two young corms of *Calla* after having grown for some time in v. d. Crone-culture-solution. They originated from a diseased plant, but were brushed with formalin before being placed in the culture-solution.

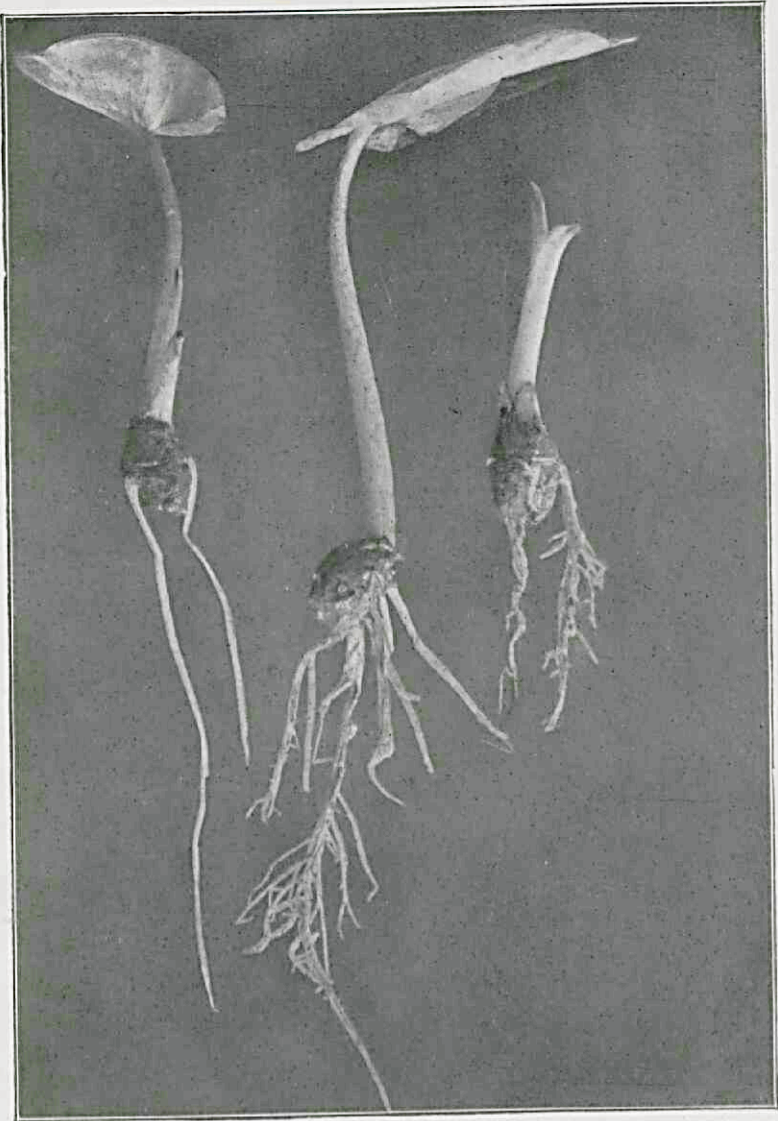


Plate 2

The two plants at the right originated from a diseased Calla and were handled in the same way as the plants in plate 1.

They were inoculated with *Phytophthora Richardiae* after the roots were well developed.

The young corm at the left originated from a healthy Calla and was also inoculated with *Phytophthora Richardiae*.

The roots have disintegrated into a glassy, afterwards slimy mass.

similar to the above described disease. He takes *Bacillus aroideae* Townsend to be the cause, but does not describe any inoculation experiments with this organism. Townsend (36) says about the symptoms of his disease: „The writer found the Calla's rotting off usually at or just below the surface of the ground, the disease sometimes extending down into the corm, sometimes upward into the leaves, and frequently in both directions. Occasionally the disease seemed to start in the edge of the leafstalk, in the flower stalk or in some underground part of the corm, though as a rule it started at the top of the corm just above, but near the surface of the ground”.

These symptoms differ in many respects from the ones I noticed in the Calla-nurseries in Holland.

Although there seems to be a great similarity between the symptoms described by Bewley, and those of the disease in Holland, I am not able to say if the two diseases are identical, as I never had the chance of examining Bewley's specimens.

Bewley used a solution of formalin diluted from 1 to 49 for control measures. He leaves the corms in this solution for an hour, without any damage being done to them. This proved to be a very good way for controlling the English disease.

In the meanwhile, the formalin treatment has been applied at one of the Calla-lily nurseries in Holland, and it was found, that the disease was far less universal than before. It is hoped, that on applying the method a second time, the disease will be entirely got rid of.

CHAPTER IV

ROT OF CUTTINGS

§ 1. General discussion

Especially in the last few years many cases are reported of cuttings which became infected by different species of *Pythium*. In 1910 Peters (24) described the „Schwarzbeinigkeit“ (Blackleg) of geranium caused by *P. de Baryanum*. In 1922 van Poeteren (30) also reports the presence of *P. de Baryanum* on geranium cuttings.

A publication on Blackleg of Pelargonium by W. Buddin and E. Wakefield (9) appeared in 1923, in which *P. de Baryanum* and *Botrytis cinerea* are named as the cause of the disease.

In 1924, H. Pape (23) reported *P. de Baryanum* as a parasite on carnation cuttings.

A detailed description of *Pythium* species occurring on geranium, was published by Braun (8) in 1925. He describes two new species, *Pythium splendens* and *P. complectens*.

In May 1925 some chrysanthemum cuttings, which had suddenly stopped growing, were sent to me to be examined. It became evident, that the roots were suffering from a rot. I soon succeeded in isolating a phycomycete from these decaying roots. This fungus grew well on oatmeal-agar; it produced no oogonia however, but a large number of conidia or swarmsporangia were formed in agar-cultures. In order to be able to identify it, it was necessary to find out in which way the zoospores were formed. I therefore placed

a tuft of the mycelium from agar-cultures in a watchglass with water. It appeared, that this is not the right method for obtaining zoospores. It was possible however to identify *this* fungus without seeing swarmspores, when it is placed in water.

If, for instance, the mycelium in the watchglass was allowed to stand for about twelve hours without the water being refreshed, numerous sporangia were formed in chains. The first formed specimen of such a chain often has a somewhat obovate shape, the others are round. This corresponds with the description of *Pythium intermedium* given by de Bary (4). He also mentions the absence of oogonia and antheridia. The chains of sporangia are so characteristic, that it is nearly impossible to mistake it for other Phycomycetes. De Bary found *Pythium intermedium* together with *Pythium de Baryanum* in dead seedlings of *Lepidium* and *Amaranthus*. Inoculation experiments however proved that it is impossible for *P. intermedium* to penetrate into these seedlings while they are living. But, according to de Bary it easily penetrates into the tissues of healthy prothallia (*Equisetum*, *Todea*, *Ceratopteris*). *P. intermedium* is also mentioned at length by Butler in his publication on *Pythium*.

He states:

„It is curious, that this species, which is common as a soil saprophyte, should have been so rarely observed. I have so regularly obtained it from water, in which were suspended fresh roots of *Abutilon*, that I at one time suspected parasitism on this plant. I however, failed to induce it to attack living roots. De Bary equally failed to infect *Lepidium sativum* and *Amaranthus* with it.

He, however, observed its parasitism on prothalli. Atkinson found, that it was responsible for a disease of fern prothalli in the United States. The affected plants were soft, limp and darker in colour than healthy ones. A high temperature, moist soil and air, and insufficient light and ventilation, are the chief predisposing factors to damping-off. When the soil is infected, it should be replaced by fresh sterilized earth".

I also came across this species several times. From Louise Solberg (Oslo) I received a strain of the same fungus, which she

had isolated from diseased tubercles of a lupin. It is therefore probable, that *P. intermedium* often occurs in vegetable mould. It is easily distinguished from the other Phycomycete-types with round sporangia and without oogonia, which are so often found on decaying roots or in the soil, by the often long chains of sporangia, which are formed when the mycelium is placed in water.

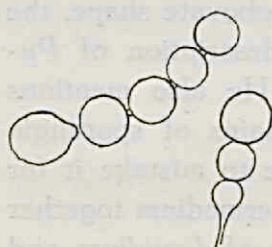


Fig. 8
Chains of sporangia or
conidia of *Pythium in-*
termedium. 366 X

The solution of Petri is again successful in obtaining zoospores.

If the mycelium is left for a few days in this solution and then transferred to a watch-glass with pure water, swarmspores appear after a few hours or sometimes after a day. If the mycelium is straightaway transferred from agar-cultures to water most of the sporangia germinate as conidia.

Inoculation experiments now had to prove, that *P. intermedium* is indeed able to infect chrysanthemum-plants and cause a retarda-

tion in growth.

When I examined the infected plants two months afterwards to study the course of the disease, it appeared that it was not of a very serious nature. All plants, also those which had been most badly infected, had outgrown the disease. This was probably also due to more favourable conditions of temperature and humidity.

Up to now the presence of *Pythium intermedium* on chrysanthemum or other cuttings has not been reported. I also wanted to inoculate geranium cuttings with it, as they are apparently easily attacked by different *Pythium* species. It is naturally thought, that the fleshy, juicy stems of geraniums are more easily attacked by Phycomycetes than the woody stems of chrysanthemums.

§ 2. Inoculation experiments

For comparison I also carried out some inoculation experiments with *Pythium de Baryanum* isolated from peas, *Pythium complectens* and *P. splendens* obtained respectively from Braun and from Leonian and with a Phycomycete corresponding with *P. de Baryanum* var. *geranii* Braun.

This last fungus is one of those forms without oogonia which appeared when diseased roots were layed out on agarplates. It is characterized by fairly large, dark, round conidia and many chlamydospores of different sizes. I did not succeed in obtaining swarmspores or oogonia from it.

The best results with inoculation-experiments were obtained when the lower ends of the cuttings were rubbed with the mycelium from agar-cultures and then placed in pots with coarse sterilized sand. In order to get as much moisture as possible, the pots were placed in socalled „sun-catchers”, small glass houses built by means of thick iron wires.

After a few days, the first symptoms of the disease appear. The base of the cutting becomes black and the colour slowly spreads towards the top. This phenomenon was also discussed by Braun. The infection by *Pythium splendens* spreads through the whole cutting, while that of *P. complectens* stops a few c.M. from the base of the slip. I think that the typical dark black discoloration of the tissues of geranium is characteristic for infection by *Phycomycetes*, while with *Botrytis*-infection a more brown discoloration is typical. It appears, that on a whole geranium-cuttings are more easily infected than chrysanthemum cuttings.

It only seldom happens that a chrysanthemum cutting is so badly infected that it is killed at once. It is probable however, that the somewhat diseased plants may show a temporary retardation in growth.

As I always used fairly young slips I have not been able to determine if the rot may also attack the roots, as one would suspect when examining the originally diseased plants, which were much older than the cuttings used by me. In any case, very special conditions of temperature and humidity must appear, before such a disease will be the result.

Inoculation experiments proved that *Pythium intermedium* is strongly pathogenous for geranium cuttings, having the same effect on them as *P. splendens*. Chrysanthemum cuttings on the other hand, are only slightly attacked by it.

Pythium de Baryanum var. *geranii*, like *P. de Baryanum* proved to be pathogenic for chrysanthemum cuttings, while *P. splendens* also attacked chrysanthemum slips.

Some results of the inoculation experiments are given below:

1. Result of inoculation experiment in a hothouse with mean temperature about 20°, on 11th. February '26.

6 geraniums and 1 chrysanthemum inoculated with *P. splendens*.

On 16th. February: 5 geraniums and 1 chrysanthemum obviously infected, 1 geranium attacked at the base (this one died afterwards).

6 geraniums and 2 chrysanthemums inoculated with *P. complectens*.

These inoculations gave no result.

6 geraniums and 11 chrysanthemums inoculated with *P. intermedium*.

On 18th. February: 1 chrysanthemum and 4 geraniums obviously infected; 1 geranium black at the base, the others healthy.

2. Result of inoculation experiment carried out in the same hothouse on 23rd. April '26.

12 cuttings of chrysanthemum and 6 cuttings of geranium inoculated with *P. intermedium*.

On 26th. April: 1 chrysanthemum beginning to show black discoloration.

On 29th. April: 4 geraniums turning black, 1 beginning to show discoloration.

The other plants remain healthy.

3. Result of inoculation experiment carried out in the open in sterilized sand on 17th. and 18th. June 1926. (See table p. 27). As *Pythium intermedium* often occurs in vegetable mould, I presume, that blackleg of geranium is not only caused by *Pythium de Baryanum*, but also by *P. intermedium*. It is however necessary to isolate the fungus from the diseased geraniums in order to be able to say to what species the blackleg is due.

	Number of inoculated plants	Heavily affected	The infection stops on some distance from the base	Healthy
<i>P. splendens</i>	6 ger.; 6 chrys.	6 ger.; 1 chrys.	2 chrys.	3 chrys.
<i>P. complectens</i>	3 ger.; 3 chrys.		3 ger.	3 chrys.
<i>P. intermedium</i>	6 ger.; 6 chrys.	6 ger.	2 chrys.	4 chrys.
<i>P. de Baryanum</i>	6 ger.; 6 chrys.	4 ger.	2 ger.; 2 chrys.	4 chrys.
<i>P. de B. var. geranii</i>	6 ger.; 6 chrys.	5 ger.	1 ger.; 2 chrys.	4 chrys.
Control	5 ger.; 5 chrys.			5 ger.; 5 chrys.

CHAPTER V

ROOT ROT OF PEAS

§ 1. *General discussion*

In the last few years a great deal has been done in America towards investigating the root rot of peas caused by *Phycomycetes*. — Especially F r e d. R. J o n e s did much in solving this problem. In 1923 (16) he drew attention to the fact, that a fungus which seemed to be a *Phycomycete*, is often found in the tissues outside the endodermis of the roots of Legumes and some other plants. He did not succeed in cultivating this organism outside the roots. When it was present in large quantities it seemed to cause a disease in the roots of *Lathyrus*, but under ordinary circumstances it did not appear to do much harm. Jones therefore regards it as a kind of mycorrhizal fungus.

Perhaps it is a *Phycomycete*, but nothing further has become known about it. F. R. Jones and his assistants however, afterwards investigated other *Phycomycetes* which occur specially on pearoots. A publication by D r e c h s l e r, *Root rot of Peas in the middle Atlantic States in 1924*, appeared in 1925. (11)

The *Phycomycetes* which he isolated from pearoots are:

1. 6—8 species of *Pythium* of which 3 are usually included in *Pythium de Baryanum*,
2. 2—3 species of *Artotrogus* (with thorny oogonia),
3. *Aphanomyces euteiches*.

This last appeared to be the most pathogenic and is very fully described by F. R. Jones and Drechsler (17). The

disease especially appears on fields on which peas have been planted a few years in succession.

Drechsler further remarks about *Pythium* and *Aphanomyces*:

„Although some isolations were made from discoloured rootlets not showing evidence of being attacked by any other fungus the most prolific source for cultures of species of *Pythium* was found in the cortical tissues of the stem and larger roots bearing the oogonia and oospores of *Aphanomyces euteiches*. In fields in which the latter fungus was common, the genus *Pythium* appeared to be found occurring more abundantly in secondary relationships than in directly parasitic ones. This condition is apparently not due to any lack of potential virulence, since nearly all of the smooth forms tried out so far, as well as one spiny form, which was derived from material collected at Hamburg, N. Y. attack cucumber fruits with great readiness, generally a fair index of a moderate degree of pathogenicity.

„The widespread occurrence of even the most aggressively parasitic species of *Pythium* in dead organic matter has long been recognized, and it is evident that the tissues killed by *Aphanomyces euteiches* provide a more congenial substratum than the living parts”.

F. R. Jones and Linford published an account called: *Pea disease survey in Wisconsin*, (18) in which the damage due to species of *Pythium* is treated in one chapter.

In this chapter he states:

„When pea plants suffering from root rot are examined in the laboratory, the species of *Pythium* long known as a destructive seedling parasite will often be found present in the diseased tissue of many or all of the plants. Inoculation with some of the cultures of *Pythium* obtained in this way has shown, that the fungus is capable of preventing germination of pea seed or of destroying many of the seedlings before they emerge from the ground, and occasionally some degree of stem and root rot is produced.

„The study of the relation of *Pythium* to root and stem rot has been greatly retarded by the fact, that the cultures obtained from peas in the field have been found to belong to several species differing somewhat in pathogenicity and frequency of occurrence and

furthermore it is not at all certain, that present methods used in making isolations from plants secure cultures of all species present. Thus a vast deal of work will be required before the relation of *Pythium* species to stem and root injury of mature plants will be fully known. It may be stated, however, that inoculations made under controlled conditions have shown very slight ability in any species studied thus far to produce either stem or root rot under usual field conditions.

„During the summer, isolations were frequently made to secure cultures of any species of *Pythium*, that might be present. Since previous experience had shown, that some species are almost as frequently found in association with roots apparently healthy as with those diseased cultures were made from both. In making cultures no sterilizing agents were applied to the surface of decaying tissue because they penetrate rapidly and destroy all *Phycomycetous* mycelium quickly. Thus mycelium adhering to the outside of roots may give rise to a culture, a difficulty which can hardly be avoided.

The results of 76 isolations are summarized in a table:

	No culture of <i>Pythium</i> obtained	1 species of <i>Pythium</i> obtained	2 species of <i>Pythium</i> obtained
Healthy pea roots	10	22	14
Decaying pea roots and stems	6	16	8

„This table corroborates previous experience that cultures of *Pythium* are obtained with approximately the same frequency from healthy as from diseased plants. It also shows that in about one-third of the isolations two species were obtained, though in such cases no two species seem to be found more frequently associated than others.

„From the observations made thus far it appears that some of these species of *Pythium* occurring abundantly in the soil may at times be responsible for the death of root ends and for some rootlet injury, but only rarely for root and stemrot as it is usually known. It is possible that almost universal invasion of the root cortex of

peas and clover by a mycorrhizal fungus renders this tissue especially accessible to these species and that it is from this superficial invasion that the fungus is most frequently obtained in culture". —

In Europe, *Pythium Sadebeckianum* is reported occurring on peas. Butler (10), however, omits this species. He remarks about it: „This species occurred as a parasite on peas in Pomerania and lupines in Hamburg, doing a considerable amount of damage in 1891. The diseased peas were examined by Wittmack, who published a brief account of the pathological conditions without, however, any diagnosis of the fungus further than the measurements of bodies which resembled the oospores of a *Pythium*. Sadebeck, who saw the specimens considered it to be a new *Pythium*, allied to *P. de Baryanum* and *P. Equiseti*. The oogonia measured 32 μ in diameter. Whether it is the same as the species generally included under *P. de Baryanum*, which has been reported attacking peas in the United States, England and elsewhere is uncertain".

These are the most important publications which have appeared about the rootrot of peas caused by *Phycomycetes*.

In order to determine whether *Aphanomyces euteiches* is also found in this country, I examined several fields on which peas were cultivated. It was done in the first part of May as the plants are often attacked early in the season. Although a lot of rain fell in the spring of 1926 and conditions were therefore favourable for the development of the disease, I only found a few plants showing signs of disease. Only on one field a few rows of plants were apparently affected.

The affected plants first became yellow or mottled and then dried out. If the plants were taken out of the soil it could be observed that the roots showed signs of decay. I collected diseased plants from different parts and examined them microscopically. In the roots I found hyphae and sometimes oogonia belonging to some kind of *Phycomycete*, but I never met the peculiar hyphae of the mycorrhizal fungus described by Jones.

In isolating the fungus from diseased roots I of course had to cope with the same difficulties as Jones.

As the soil is full of *Phycomycetes* and the roots cannot be

sterilized, one never knows if an isolated species is actually the same one as that of which oogonia appear in the roots. Only inoculation experiments could prove if a certain isolated species would be pathogenic or not.

From most of the isolations a species was obtained with oogonia often possessing protuberances. I had met this fungus before in a young lupin which had shown the typical symptoms of damping-off.

When a colleague tried to isolate *Cladosporium cucumerinum* from cucumber seeds which failed to germinate the same species appeared in the isolations. Apparently it is a common soil fungus. Except this fungus with irregular oogonia, which I usually obtained from diseased roots, other forms appeared without oogonia and which were therefore hard to identify. I have however chiefly confined myself to the fungus with irregular oogonia which I shall afterwards describe as *Pythium irregulare* n. sp.

Besides I carried out some inoculation experiments on peas with one of the fungi without oogonia, which corresponds with *P. de Baryanum* var. *geranii* Braun. Although this fungus had not been isolated from peas it had appeared to be very virulent during experiments on cuttings. For comparison I also carried out some inoculation experiments with *Pythium de Baryanum* and *Aphanomyces euteiches*.

The former I had isolated from diseased pea roots, the latter was obtained as a pure culture from the United States (Drechsler).

§ 2. Inoculation experiments

For my first inoculation experiments I used humus, through which I mixed agarcultures of the fungi before sowing the peas in it. At the time I had not yet isolated *Pythium irregulare* from pea roots, so that I mainly used *Pythium de Baryanum* and *P. de Baryanum* var. *geranii*. It soon became evident that this method is not very satisfactory. The little radical was immediately attacked by the fungus and the peas did not come up. If they did, they either were very small and miserable or they outgrew the disease, but in neither case did they show the typical symptoms of root rot. Another drawback was that I had to use peas with a high percentage of germination, as it would otherwise be impos-

sible to carry out control experiments. „Wonder van Amerika”, the variety I used, had a very high germinative power, but on the other hand it is more resistant to disease than other varieties of green peas (doperwten). When working with less virulent parasites there would therefore be more chance of the inoculation succeeding with less resistant varieties of green peas, than with the Wonder van Amerika. I arranged the inoculation experiments in the following way: the peas were sown in pots containing sterilized vegetable mould mixed with oatmeal-agar cultures of the different fungi in question.

The results of one of these experiments were:

	Number of seeds	Number of plants living after 11 days
Inoculated with:		
Pythium de Baryanum (isolated from pea-roots)	15	0
Pythium de Baryanum (strain from Hartley)	15	1
Pythium de Baryanum (Strain from diseased corm of Gloxinia)	6	1 poor specimen
Pythium de Baryanum var. geranii	15	3 (one of which very small)
Control	5	5

It therefore appeared that this method is not the right one.

These difficulties could be removed by using watercultures. One then also has a better survey of the progress of the rot. Other soil fungi which in pot experiments always follow on the Phycomycetes were eliminated in this way. The roots which have grown in watercultures are clean and therefore very suitable for microscopical examination.

The seeds were allowed to germinate and the best seedlings were then sorted out and used for the waterculture-experiments.

For these experiments I used a green pea called Prins Albert.

It possesses long, thin roots, while seedlings of Wonder van Amerika have a short firm root system.

I proceeded as follows: The seeds were first sterilized with alcohol-sublimate (1 alcohol 96 % : 1 sublimate 0,1 % to prevent infection in the germinating dishes as far as possible.

They were then laid in dishes on wet filterpaper and when the roots were a few c.M. long they were used for the watercultures.

The watercultures were arranged as follows:

In the lid of a jampot three holes were made and in these holes two pieces of cork, somewhat hollowed out in the middle, were fitted. The roots of the seedlings were pushed through the round hole, which was formed between the two pieces of cork, while the cotyledons rested on the top of them.

The jampots were filled with the culture solution of v. d. Crone.

After the roots had grown in the solution for a little while, I placed a tuft of the mycelium from the fungus with which I wanted to inoculate, against them. It may of course be objected against this method that the fungus has an advantage over the plant in this way. A pea is not a waterplant and is therefore not cultivated under normal circumstances, when growing in watercultures.

Phycomycetes on the other hand are water-fungi and easily produce swarmspores in mineral solutions, so that in watercultures the circumstances are very favourable for infection. The pea seedlings, however, grow so well in v. d. Crone's culture solution, and produce such a healthy root system, that according to my opinion, they may well be compared with plants grown under normal circumstances. At the utmost one may get an exaggerated idea of the pathogenicity of the parasite, when one relies on the damage done by it to the root system of pea seedlings in watercultures.

But then the damage done to the green parts of the plant is usually less striking in watercultures than on the fields. On the fields the root rot is characterized by the drying out of the infected plants and this of course does not soon happen in watercultures: as long as a few roots remain intact the plant will not soon dry out. Although the whole aspect of the disease is therefore somewhat different in watercultures than in the pots, one may still draw conclusions about the pathogenicity of a special Phycomycete for the roots of peas. *Aphanomyces euteiches* appeared

to be the most virulent of the four Phycomycetes (*Aphanomyces euteiches*, *Pythium de Baryanum*, *P. de Baryanum* var. *geranii* and *Pythium irregulare*), with which inoculation experiments were done.

A little while after the pea roots had been inoculated with *Aphanomyces euteiches* the whole root system became limp.

They did not become as glassy and transparent as with *Pythium* inoculations but remained white. The infection is very acute. Even in watercultures the plants may shrivel up and die.

When inoculated with *Pythium irregulare* the roots were in many cases also soon badly infected. On the upper part of the plants nothing could be seen however. The growth was normal and the leaves were dark green. The roots were glassy, transparent and brown in colour, while there were only a few small side roots at the ends. These side roots may, when they are also infected, show a reddish brown ring at the place where they are attached to the main roots. When roots infected by *Aphanomyces euteiches* or *Pythium irregulare* are examined microscopically, large numbers of oogonia are found in them.

Pythium de Baryanum and *P. de Baryanum* var. *geranii* also attack pea roots in watercultures. In their method of infection they resemble *Pythium irregulare* and not *Aphanomyces euteiches*.

The plants in the control experiments remained healthy and possessed a clean white, well branched root system.

From these results I think I may conclude that *Pythium irregulare* is the cause of a root rot of peas, and that a large number of the diseased plants with decaying roots, which I found on the fields, is due to it. Finally it may be pointed out, that it is hard to distinguish *Pythium irregulare* from *Pythium de Baryanum* when the infected roots are examined microscopically.

The protuberances of the oogonia are very hard to find in the tissues of the roots.

In culture however they are easily distinguished.

§ 3. *Description of Pythium irregulare nov. spec. in culture*

Many of the isolations which I made of this *Pythium* showed different variations. I specially kept three of them in culture, one

from peas, one from lupins, one from cucumber seeds. Macroscopically these fungi look exactly like other *Pythiums*. I usually cultivated them on oatmeal-agar on which they made the usual white woolly growth of mycelium. I tried to get swarmspores by the method of Petri. The strain isolated from cucumber seeds produces a few swarmspores in this way, but there were too few to observe the discharge of the swarmspores from the sporangia, or to be able to count the cilia. For the purpose of counting the cilia, I placed a drop of water with swarmspores on a slide with a drop of eosine-sublimate (following the advice of Mr. v. Luyk).

The swarmspores were immediately fixated and coloured by the eosine before they withdrew the cilia. It is obvious that one can only apply this method when there are a fairly large number of swarmspores at one's disposal. It was impossible for me to count the cilia, as the strain from cucumber only produced a few, the strain from lupin only very few single ones, and the strain from pea no swarmspores at all.

As I could not determine the way in which the swarmspores were formed the difficulty was to

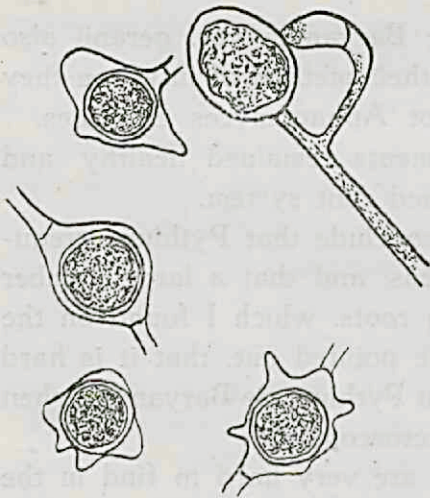


Fig. 9

Oogonia of *Pythium irregulare* isolated from cucumber-seed. 550 \times

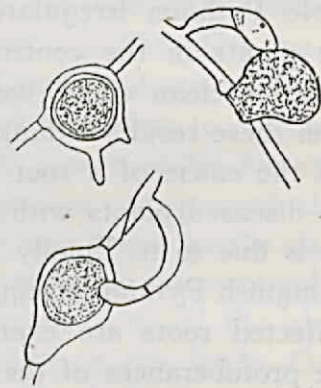


Fig. 10

Oogonia of *P. irregulare* isolated from pea-roots. 550 \times

decide whether the species belonged to *Pythium* or to *Phytophthora*.

I did not hesitate for one moment, however, in placing it with

the genus *Pythium*, owing to the great similarity which exists between this fungus and *P. de Baryanum*.

The only difference which really exists between this species and *P. de Baryanum* is the irregular shape of the oogonia. The oogonia often (but not always) carry protuberances, varying in size and

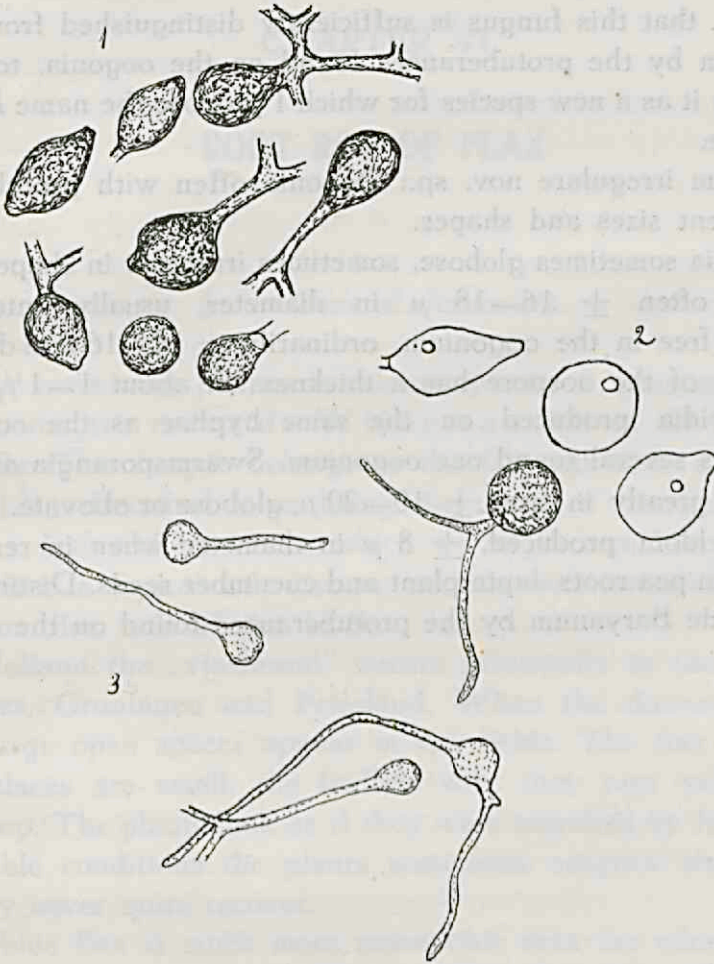


Fig. 11

- 1 Conidia of *Pythium irregulare* isolated from lupin. 550 \times
- 2 Empty sporangia and conidium of *P. irregulare* isolated from cucumber-seed. 550 \times
- 3 Germinating zoospores of *P. irregulare* isolated from cucumber-seed. 550 \times

shape, and sometimes separated from the oogonium by a transverse wall. These protuberances are especially found in cultures

a few weeks old. The oogonia are mostly intercalary and the oospores, sometimes two in number, lie free in the oogonium. The antheridia are usually of the same type as those of *P. de Baryanum*, but I never saw hypogynal antheridia. One oogonium is sometimes surrounded by more antheridia. The conidia or swarmsporangia are globose or somewhat obovate, and differ in size and shape.

I think that this fungus is sufficiently distinguished from *P. de Baryanum* by the protuberances found on the oogonia, to justify recording it as a new species for which I propose the name *Pythium irregulare*.

Pythium irregulare nov. sp.: Oogonia often with protuberances of different sizes and shapes.

Oogonia sometimes globose, sometimes irregular in shape. When globose, often $\pm 16-18 \mu$ in diameter, usually intercalary. Oospore free in the oogonium, ordinarily $\pm 14-16 \mu$ in diameter. The wall of the oospore has a thickness of about $1-1\frac{1}{2} \mu$.

Antheridia produced on the same hyphae as the oogonium, sometimes several round one oogonium. Swarmsporangia and conidia vary greatly in size, $\pm 10-20 \mu$, globose or obovate. Swarmspores seldom produced, $\pm 8 \mu$ in diameter when in rest. Isolated from pea roots, lupin plant and cucumber seeds. Distinguished from *P. de Baryanum* by the protuberances found on the oogonia.

CHAPTER VI

ROOT ROT OF FLAX

§ 1. General discussion

When examining pea roots microscopically I sometimes observed *Asterocystis radialis* de Wildeman (37) in the parenchymal cells. This fungus belongs to the Chytridiaceae and Marchal (21) described it as being the cause of the Belgian and Dutch „vlasbrand”, a name, which we might translate by „flax-fire”. It is a typical wilt-disease, not identical however with the *Fusarium* flax wilt of the U. S.

In Holland the „vlasbrand” occurs principally in the northern provinces, Groningen and Friesland. When the disease is very acute large open spaces appear in the fields. The flax plants in these places are small, the leaflets wilt, they turn yellow and shrivel up. The plants look as if they were scorched by fire. Under favourable conditions the plants sometimes outgrow the disease, but they never quite recover.

The blue flax is much more susceptible than the white variety. In severe epidemics however, as in 1926, the white flax is also attacked. On old heavy clay soils, the „brand” rarely occurs, but on the lighter sandy clay, the disease is very common. It is remarkable that even in the new „polders”, where flax is grown for the first time, the „brand” may occur. This disease is of a severe nature; the yield on those fields is extremely small. Dr. Zijlstra, chief of the botanical department of the Groningen Experiment Station, called it „a pest for our best flax regions”.

Marchal found *Asterocystis radialis* in all the „brand” specimens from different parts of Belgium. He proved the parasitic nature of this fungus by carrying out inoculation experiments. He grew flax in watercultures and inoculated the nutrient solution with *Asterocystis*; the result was a thorough infection of the roots. By growing the plants in infected soil, he also obtained diseased plants. The presence of *Asterocystis* in the roots is easily ascertained by a simple staining process with potassium-iodide. Marchal then determined that the chlamydospores and swarm-sporangia were stained a deep brown-red. Where the symptoms of the disease were concerned Marchal remarks: „The less water the plants have at their disposal, the stronger they react on the lack of water, which becomes evident after infection and the more striking are the symptoms of the „brand” ”.

The moisture, which by influencing the dissemination of the swarmspores helps to spread the disease, on the other hand diminishes its pernicious influence.

Marchal's investigations are the only ones published about this disease. Dr. Zijlstra informed me how to obtain infected plants in spring. He therefore sent me a sample from „brand” soil, which had been collected two and a half years ago. He advised me to grow flax in nutrient solutions (van der Crone). The bottom of the jar was covered with a thin layer of the soil. This infection succeeded beautifully. A large part of the root-cells was filled up with the *Asterocystis* sporangia. The plants of course, having their roots in the liquid, did not shrivel like those in the field, although they had a sickly appearance. The control plants were quite healthy.

When growing peas in the same solution, I also noticed a slight infection of the roots. In this case however the plants remained quite healthy.

In the summer of 1926 I visited the flax district of Groningen and collected plants of a „brand” spot in the field. I was there told by one of the foremost farmers, that these plants were typical „brand” plants.

The examination of these specimens however revealed relatively few sporangia of *Asterocystis*, but a great many round spiny bodies, which struck me as being oogonia of a Phycomycete. Later

in the summer I had the opportunity of examining flax roots from Friesland, and again I noticed the same phenomenon; the spores of *Asterocystis* however were more abundant in the roots here.

The question now arose, whether the root rot of flax might be caused by the *Phycomycete* in question, as well as by *Asterocystis radialis*. It would not be surprising if the wilting and shriveling of the plants were caused by different parasites. The „vlasbrand" would then be a complex of diseases.

It is possible that several kinds of root parasites might nearly have the same effect. We know f. i. that *Thielavia basicola* may cause a rootrot of flax. —

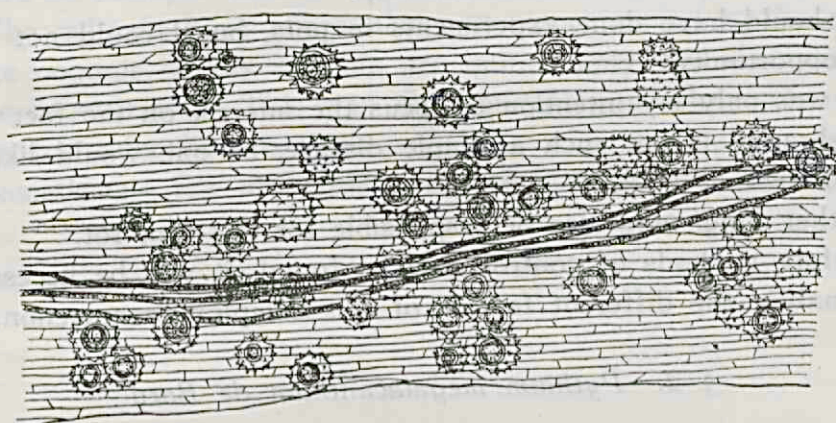


Fig. 12

Part of flax-root crushed under the cover-slip after inoculation with *P. megalacanthum*. $\pm 70 \times$

The *Phycomycete* proved to be *Pythium megalacanthum* de Bary (4), which was isolated very easily from the decaying roots. I thus intended to make inoculation experiments on the roots, with the exclusion of *Asterocystis*. For this purpose the flax seeds were germinated on wet filter-paper. The young plants were put in the nutrient solution and mycelium from the pure cultures on oatmeal-agar was smeared on the rootlets.

During the first days after the inoculation the roots apparently develop normally. After some days however, the top of the principal root begins to soften. This „rotten" end is separated from the healthy part by a red ring.

As the rotting process advances, the ring is slowly replaced.

Except the main root, the lateral roots are also attacked. All these infected roots become glassy and limp.

Fig. 12 shows a piece of flax root from such an artificially infected waterculture. The spiny oogonia appear in large numbers in the roots.

It therefore appears that it is possible to cause a rot of the flax roots by *Pythium megalacanthum*, even when *Asterocystis radialis* is absent.

As Marchal already noticed with his experiments with *Asterocystis*, the plants in watercultures do not suffer very much from wilting. This also counts for *Pythium megalacanthum*. The aerial parts of the plants do not shrivel up.

I should have done experiments in pots, but I could not find the opportunity.

It was only my intention to focus the interest on the fact that „vlasbrand” is not such a simple disease, as one would like to imagine.

Other fungi may also be responsible for this root rot.

Before methods of control may be found, it will be necessary to analyse the different factors of this complex-phenomenon.

§ 2. *Pythium megalacanthum* de Bary

The large spiny oogonia of the Phycomycete which I found in flax roots, naturally reminded me of the illustrations of the oogonia of *Pythium megalacanthum* given by de Bary. De Bary found *Pythium megalacanthum* in seedlings of garden-cress which had already been attacked by *Pythium de Baryanum*. He tried to infect the seedlings by inoculating them with *Pythium megalacanthum* but did not succeed. He then inoculated prothallia of the fern *Todea africana* with it and some of them were attacked. He always used zoospores for his inoculations. According to Butler this fungus has never been described again except on one occasion when Schröter states, that he believed he found it in the stems of *Veronica hederifolia*.

The Phycomycete which I isolated from flax seemed to correspond very well with the description given by de Bary for *P. megalacanthum*. The oogonia were often larger in size than the

measurements given by de Bary, but they vary so much that this is of no importance. De Bary already pointed out that the spores of *P. megalacanthum* vary more in size than those of *P. intermedium*. The drawing of the infected flax root (fig. 12) shows the variations in size of the oogonia, which range from ± 30 to 70μ . I did not succeed in getting zoosporangia. Amongst his different cultures on seedlings of garden-cress, de Bary also had one form which possessed numerous oogonia and only few swarmsporangia. Unfortunately I only possess one isolation and perhaps that is the reason for the absence of the swarmsporangia. Even the method of Petri gave no results. *P. megalacanthum* grows on oatmeal- and on cornmeal-agar but not on cherry-agar.

The cultures have a granular appearance owing to the many large oogonia. Many oogonia are quite empty and the contents have degenerated. This was also noticed by de Bary who takes them for unfertilized oogonia.

Sometimes a few antheridia are present round one oogonium.

CHAPTER VII

MYCOLOGICAL ANALYSIS OF THE VEGETABLE MOULD

In connection with the inoculation experiments, it was necessary to know whether many Phycomycetes appear in vegetable mould in which the various diseased plants, which I had examined, grew. If many Phycomycetes occurred, the experiments with unsterilized soil would be worthless. With my first experiment in which the peas were sown in unsterilized soil, I already noticed that an infection by *Pythium de Baryanum* appeared in the control experiment. Unsterilized soil could therefore not be used.

When a decaying root is laid in a petridish on cherry- or on oatmeal-agar several *Pythiums* often start growing out of it. In the chapter about the root rot of peas I already pointed out that according to Drechsler (11) this may happen even if the rot is caused by a different parasite (in that case by *Aphanomyces euteiches*). F. R. Jones and Linford (18) even isolated several *Pythiums* from *healthy* pea roots.

One may gain an impression of the great number of Phycomycetes occurring in vegetable mould by laying out roots of different plants.

The many types which appear, make a research about Phycomycetes rather difficult at first.

One comes across several forms without oogonia or without swarmsporangia. With such isolations one often meets *Pythium de Baryanum* and the form called *P. de Baryanum* var. *geranii* by Braun. These forms were also often isolated from various seedlings.

One of my colleagues tried to isolate *Cladosporium cucumer-*

inum from cucumber seeds which failed to germinate in vegetable mould, but on one occasion he obtained different *Pythiums* instead. He handed them over to me and it appeared that *Pythium intermedium* and *Pythium irregulare* were present. From these seeds I also isolated a form with dark, somewhat spiny oogonia. I afterwards never came across this form again. Although the seeds were put to germinate in a hothouse, the temperature could hardly be the cause of the occurrence of this form, as many of my own seedlings were also cultivated at about the same temperature.

From decaying roots of pansies which had been grown in another soil-sample at a lower temperature, I isolated an *Aphanomyces* which looked very similar to *Aphanomyces euteiches*, but which, in my infection experiments, did not attack pea roots. Unluckily I did not have time to carry out inoculation experiments with it on pansies, and I have therefore not been able to ascertain whether this *Aphanomyces* is the cause of the acute root rot of pansies appearing in our experimental garden. From Calla-lily-roots I isolated, besides *Phytophthora Richardiae*, a few other forms, probably also *Pythiums*.

I never obtained *Phytophthora* species in the same way and apparently they do not lead the same saprophytic life in the soil as the different species of *Pythium*. The question of the temperature, which may be of importance here, was not investigated however. —

It is evident that vegetable mould is full of different *Phycomycetes* and that inoculation experiments with unsterilized humus would not be trustworthy.

It would lead me too far to discuss the different forms in this publication.

I also made some isolations from clay-soil. The difficulty with these isolations was that the dishes were often overgrown by a *Mucor*. But here again other forms than those which I had isolated from vegetable mould appeared.

SUMMARY

1. It has become evident that there are no sound reasons for regarding *Pythiomorpha* Petersen and *Blepharospora* Petri as constituting new genera. There is no objection against including them in the genus *Phytophthora*.

In practice it is not desirable that the genera *Pythium* and *Phytophthora* should be placed in one genus, although previous investigators have shown that theoretically there is no difference between them.

2. Inoculation experiments proved that a serious root rot of Calla-lilies, which causes a considerable damage in the nurseries, is caused by a *Phytophthora* species. As this species is not identical with any of the ones previously described, the name *Phytophthora Richardiae* n. sp. is proposed.

A treatment of the Calla corms with formalin, which has already yielded good results in practice, is given as a control measure.

3. It appeared that *Pythium intermedium* de Bary, isolated from chrysanthemum cuttings, which had stopped growing, is strongly pathogenic for geranium cuttings and may also attack chrysanthemum cuttings. Inoculation experiments were also undertaken with geranium cuttings, as a publication by Braun about the rot of these slips appeared in the United States.

For comparison I also inoculated with the species *P. splendens* and *P. complectens* named by Braun. These species, which had been kept in culture at the Centraal Bureau voor Schimmelcultures for some time, were still very virulent. *P. splendens* also attacked chrysanthemum cuttings. Inocula-

tion experiments with the strains of *P. de Baryanum* and *P. de Baryanum* var. *geranii* which I isolated, gave positive results with geranium and chrysanthemum cuttings.

4. A *Pythium* with irregular oogonia sometimes with protuberances, was isolated from the roots of a large number of the pea plants I found suffering from root rot.

As far as I know this species has not been described before and I therefore give the description and propose the name *Pythium irregulare* n. sp.

Inoculation experiments in watercultures proved that it attacks the roots of peas and is the cause of a root rot. I never came across *Aphanomyces euteiches* in peas, but inoculation experiments showed that it is more virulent than *Pythium irregulare*.

5. Typical „vlasbrand” has been ascribed to *Asterocystis radialis* de Wildeman.

The symptoms of „vlasbrand” of course occur whenever something is the matter with the roots. When examining the diseased plants I often found *Pythium megalacanthum* de Bary present in the roots. Inoculation experiments showed that this species attacks the roots of flax. As I found this species in the roots of diseased plants from different districts I think I may conclude that like *Asterocystis radialis* it may be one of the causes of the disease. A thorough re-investigation of vlasbrand is however necessary.

6. Inoculation experiments with *Phycomycetes* on roots are very much simplified by using watercultures.
7. It appeared that various *Phycomycetes* may be isolated from decaying parts of plants grown in vegetable mould. These forms are often very hard to identify owing to absence of oogonia or sporangia.

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STELLINGEN

I.

De genera *Blepharospora* Petri en *Pythiomorpha* Petersen moeten bij het geslacht *Phytophthora* de Bary worden gevoegd.

II.

De meerdere of mindere resistentie van vlasrassen tegen infectie door *Fusarium lini* berust op het voorkomen bij die rassen van een grootere of kleinere hoeveelheid glucoside.

III.

De halophyten van het Noord- en Oostzeestrand zijn niet xerophytisch aangepast.

IV.

De pseudo-vacuole der blauwwieren, die voornamelijk optreedt bij soorten, welke waterbloei veroorzaken, heeft een protoplasmatischen inhoud.

V.

De verdeeling der water-organismen in saprophielen, saprotoleranten en obligaat katharoben verdient de voorkeur boven die in saproben en katharoben.

VI.

Het chlorophyl is in de levende cel aanwezig opgelost in een lipoid.

VII.

De *Laboulbeniaceae* zijn op te vatten als kleurlooze *Florideae*.

VIII.

De droogmaking van de Zuiderzee zal ten gevolge hebben, dat Noord-Holland ten Noorden van het Noordzee-kanaal weer bevolkt wordt met typische zoetwater-organismen, die er thans niet voorkomen.

