Title: First report of the sexual stage of the flax pathogen Mycosphaerella linicola in France

and its impact on pasmo epidemiology

Authors: Delphine Paumier ^{1*}, Blandine Bammé ^{2*}, Annette Penaud ², Romain Valade ^{1**},

Frédéric Suffert 3**

Addresses:

¹ Laboratoire de Pathologie Végétale, ARVALIS Institut du Végétal, 78850 Thiverval-Grignon,

France

² Terres Inovia, 78850 Thiverval-Grignon, France

³ Université Paris-Saclay, INRAE, AgroParisTech, UMR BIOGER, 78850 Thiverval-Grignon,

France

*/** equal contributions

corresponding author: frederic.suffert@inrae.fr, ORCID 0000-0001-6969-3878

Abstract

We performed a three-year field survey in France to characterize the dynamics of sexual

reproduction in Mycosphaerella linicola, the causal agent of pasmo, during the interepidemic

period. Cohorts of fruiting bodies were sampled from linseed straw during the autumn and

winter and carefully observed, focusing on pseudothecia, asci and ascospores. A sequence of

experimental steps corresponding to Koch's postulates confirmed in July 2014, for the first time

in France and continental Europe, the widespread presence of the sexual stage of M. linicola in

plant host tissues. The developmental dynamics of pseudothecia on straw, expressed as the

change over time in the percentage of mature pseudothecia, was similar in all three years.

Pseudothecia appeared in late summer, with peak maturity reached in October. A temporal shift,

thought to be due to early autumn rainfall, was highlighted in one of the three years. These

observations suggest that sexual reproduction plays a significant role in the epidemiology of

pasmo in France. A resurgence of M. linicola infections in spring flax is thought to have

occurred in recent years, due to the increase in the area under flax. The presence of the sexual

stage of this pathogen probably increased the quantitative impact of residues of winter linseed

(used for oil) and flax straw (left on the soil for retting and used for fibers) as an interepidemic

'brown bridge'. This case study highlights how certain parts of a disease cycle, in this case the

sexual phase, can become crucial due to changes in production conditions.

Keywords

Flax, fungal pathogen, linseed, primary inoculum, pseudothecia, Septoria linicola, sexual

reproduction

Introduction

Mycosphaerella linicola Naumov, anamoph Septoria linicola (Speg.) Garassini, is the causal

agent of pasmo, a disease affecting both 'flax' and 'linseed' (Linum usitatissimum L.), crops

grown for fiber and oil production, respectively. This plant-pathogenic ascomycete affects

production in many flax-growing areas around the world, including Europe. Pasmo was first

detected in Argentina in 1909 (Spegazzini, 1911). In Europe, it was found in Yugoslavia in

1936 (Rost, 1937) and in Ireland in 1946 (Loughnane et al., 1946). More recently, this disease

has caused significant yield losses in South Dakota (Ferguson et al., 1987) and in England, on

winter linseed (Perryman & Fitt, 1999).

Pasmo usually starts on the lower leaves of young plants. During the growing season of the plant, *M. linicola* is propagated clonally by splash-dispersed asexual pycnidiospores (filiform 1-3 septate, straight or slightly curved conidia; Sivaneson & Holliday, 1981), leading to an upward progression of the disease on stem and leaves. Pasmo affects all the aboveground parts of the flax plant, causing leaf spots, a loss of flowers and capsules, and weakened pedicels (Fergusson et al., 1987). The disease also causes elongated brown lesions, which coalesce to produce mottled bands that encircle the stem (Colhoun & Muskett, 1943; Perryman et al., 2009). This 'girdling' effect and the premature death of the plant has an impact on yield and fiber quality, and may be mistaken for premature ripening (Sackston & Carson, 1951; Perryman et al., 2000).

M. linicola has been reported to be seed-borne and to survive on crop residues, like many other pathogenic ascomycetes. The sexual stage was initially reported to be absent in North Dakota (Brentzel, 1926), but several plant pathologists identified structures found on flax stubble as sexual fructifications, in Argentina (Wollenweber, 1938) and Germany (Kruger, 1941). Sackston (1949) identified structures that he believed to be 'pseudothecia' in Manitoba (Canada), but he was unable to complete Koch's postulates. Despite partial evidence for a sexual stage being repeatedly reported, this pathogen has long been thought to survive the interepidemic period on infected flax straw as a saprotroph (mycelium and old pycnidia; Gillis, 2009), with dispersal purely by rain splash or insects during the growing season (Christenson, 1952; Sackston, 1970), and long-distance carriage only in or on seeds. Sanderson (1963) was the first to identify and describe asci and ascospores from wild flax (Linorum marginale) in New Zealand, but, unfortunately, was unable to establish cultures. Pseudothecia resemble black spherical dots, 75-120 μm across, whereas asci are oblong, bitunicate, eight-spored and 30-50 × 8-9 μm in size, and ascospores are fusiform, hyaline two-celled, constricted at the septum,

and 13-17 × 2.5-4.0 μm in size (Sivaneson & Holliday, 1981; Vakhrusheva, 1986). It was not

until Perryman et al. (2009) collected air-borne ascospores thought to be those of M. linicola

that a significant role of the sexual stage in the pasmo epidemiology could be affirmed. These

authors proposed a pathogen life cycle including a sexual phase during the winter, in which the

pathogen survives as pseudothecia, although these fruiting bodies were not observed in their

study. An important role of the sexual stage and the recombination resulting from it is consistent

with the high levels of genetic diversity demonstrated for two populations from Manitoba

(Grant, 2008).

Pasmo incidence and severity have increased over the last two decades in France (no attack

reported in French trials in 1988 and 1990; Fitt et al., 1991), in parallel with the increase in the

area under Linum usitatissimum L., particularly in the form of flax for linen production (50,000

ha in the 1990s vs. 120,000 ha in 2020; data CIPALIN). During this period, France has become

the world's largest producer of scutched flax fibers (580,000 t in 2017, i.e. about 75% of the

world's production; FAOSTAT). Nevertheless, the sexual stage of M. linicola – which probably

plays a key role in the epidemiology of this disease – has never before been described in France,

or elsewhere in continental Europe.

In this study, we aimed to characterize the sexual reproduction dynamics of M. linicola on flax

straw during the interepidemic period, to assess its role in the early stages of pasmo epidemics

in French production conditions.

Materials and methods

Description of the different forms of M. linicola during the intra- and interepidemic periods

- Pasmo symptoms and both the asexual and sexual stages of M. linicola were observed on flax
stems, leaves and straw collected from two fields in 2015-16 at the INRAE-Terre Inovia
experimental station (Thiverval-Grignon, France). Fruiting bodies were collected with a needle,
crushed and examined under a microscope, after mounting in methylene blue solution. Pycnidia
and pycnidiospores (Figure 1A-B) were obtained from typical lesions on living flax leaves.
Pseudothecia, asci and ascospores were identified on pieces of flax straw left to dry outside
(Figure 1C-G).

Sequence of experimental steps demonstrating the presence of the sexual form – A sequence of experimental steps corresponding to classical Koch's postulates was developed to confirm that the observed fruiting bodies did indeed correspond to the sexual stage of M. linicola and could potentially play a role in pasmo development (Figure 2). Flax stem residues bearing putative M. linicola pseudothecia were identified (①). We used a technique for ascospore collection derived from that developed by Suffert & Sache (2011) to study the sexual stage of Zymoseptoria tritici on wheat residues to isolate them. Dry fragments of flax straw were spread on wet paper in a moist box (24 × 36 cm). Petri dishes containing PDA medium (potato dextrose agar, 39 g L⁻¹) were placed upside-down above the fragments (②). The box was placed at 18°C in the dark for 12 h. The Petri dishes were closed and incubated in the same conditions. Three days later, white yeast-like colonies similar to those obtained by Wollenweber (1938) from S. linicola pycnidiospores appeared (3). A conidial suspension was prepared by flooding the surface of a 3-day-old culture of a single-spore colonies obtained after one round of singlespore isolation with sterile water and scraping the surface of the agar with a glass rod (4). Leaves of a flax plant grown in a greenhouse were inoculated by applying this conidial suspension to the adaxial surface, with a paintbrush (⑤). The plant was enclosed in a transparent polyethylene bag containing a small amount of water to maintain humidity levels and, thus,

promote infection. Pycnidia and cirrhi, accompanied by symptoms, appeared on the inoculated leaves 10-13 days after inoculation (©). Cirrhi were collected with a needle (⑦) and deposited on PDA medium in a Petri dish for reisolation of the pathogen in its yeast-like form (®). Clusters of several hundred pycnidiospores released from other cirrhi mounted in methylene blue solution were examined under the microscope. At the same time, putative pseudothecia were observed on flax straw collected in the field and examined in a similar manner (9). Dynamics of M. linicola on flax straw during the interepidemic period in French conditions - During three growing seasons (2014-15, 2015-16, 2016-17) straw was collected from flax naturally infected with M. linicola in a field planted with a winter linseed varietal mixture (cv. Blizzard, Sideral, Cristalin, and Angora). The straw was collected just after harvest in July and stored outdoors, on grass, at the INRAE-Terre Inovia experimental station. In 2014-15, the straw was moistened before storage, whereas, in 2015-16 and 2016-17, it was left on the ground directly, without treatment. The straw was examined once weekly from 30 October 2014 to 26 March 2015 for the 2014-15 season, from 27 August 2015 to 6 April 2016 for the 2015-16 season, and from 21 July 2016 to 9 March 2017 for the 2016-17 season. An automatic weather station located 200 m from the pile of straw recorded hourly rainfall and air temperature at a height of 2 m. Each week, we observed a sample of 20 straws under a binocular microscope, to check for the presence of M. linicola pycnidia and pseudothecia. As soon as these structures appeared, a subsample of five infected straws was incubated in a moist box for 24 h. Five fruiting bodies were selected from each straw (25 in total) and dissected under a binocular microscope according to the protocol developed by Poisson (1997) for Leptosphaeria maculans. Fruiting bodies were placed in a drop of water on a glass slide and crushed with two needles to release their contents. A drop of methylene blue solution was added and the slide was covered with a coverslip. The preparation was observed under a microscope (magnification

×400), to distinguish pycnidia from pseudothecia. The M. linicola pseudothecia were classified

on the basis of developmental stage, taking into account the degree of maturity of asci and ascospores, as proposed by Toscano-Underwood et al. (2003) for *L. maculans* and *L. biglobosa*. Four classes were defined and used: (i) 'immature' pseudothecium, with no asci or ascospores; (ii) 'maturing' pseudothecium, with differentiated asci but no ascospores; (iii) 'mature' pseudothecium, with at least one ascus containing eight differentiated ascospores; (iv) 'empty' pseudothecium, from which all the ascospores had been discharged (**Figure Suppl. 1**). A pseudothecium was considered to have reached one of the four developmental classes when the first asci/ascospores in the pseudothecium had reached the stage of development considered.

Results

M. linicola pseudothecia and ascospore-containing asci were observed in July 2014 for the first time in France, and are described here (Figure 1 and 3). Koch' postulates were completed and showed that ascospore-derived strains were able to generate typical pasmo symptoms on flax leaves after inoculation with a paintbrush and exposure to high-moisture conditions (Figure 2). Based on weekly counts of M. linicola fruiting bodies (pycnidia and pseudothecia) on samples of 20 infected straws, and the dissection of five pieces of straw under a binocular microscope, we were able to determine the dynamics of pseudothecium formation and maturation over the interepidemic period, from August to April, in three successive growing seasons (2014-15, 2015-16, 2016-17) (Figure 4). This epidemiological survey provides the first evidence for the widespread presence of the sexual stage of M. linicola in France just before the emergence of the winter flax crop. The highest proportion of mature pseudothecia, ranging from 60% to 100%, was recorded in October, regardless of the year considered. The dynamics of pseudothecium maturation were similar in 2014-15 and 2015-16, with a peak in early October, followed by a steady decrease to below 20% after December. The peak was delayed by one

month (early November) in 2016-17. The 2014-15 survey did not begin until early November, and we were, therefore, unfortunately, unable to determine when peak pseudothecium maturity occurred. The first symptoms of pasmo were detected on flax seedlings (cotyledons) in the field on 25 November, 2015 (second growing season) and on 14 December, 2016 (third growing season). A similar assessment was performed in the first growing season, but at a later stage, making it impossible to determine whether symptoms occurred as early in the autumn as in subsequent seasons. The identification of *M. linicola* was confirmed by a TaqMan qPCR assay. The intron sequence of the EF1-α gene was amplified using the specific primers 'SeptoUP' (5'-TTGCCCCTCCAATTCTGGTG-3'), 'SeptoLOW' (5'- ATGTGTTAAAAGTGTTGTGTGC-3'), and the TaqMan probe 'Sonde Septo' (5'-FAM-CGAGAATTTTGGGCTTTTGCGGCTC-BHQ1-3'). The specificity of primers was confirmed with DNA extracted from pure cultures of 50 strains of M. linicola and of 24 other fungal species, including the wheat pathogen Z. tritici and the flax pathogens Sclerotinia sclerotiorum, Rhizoctonia sp., Verticillium dahliae, Pythium sp., Phoma exigua var. linicola and Kabatiella lini.

The year in which symptoms were detected earliest on emerging plants (late November 2015 vs. mid-December 2016) was also the year in which pseudothecium levels peaked earliest (mid-September 2015 vs. late-October 2016). This suggests that early symptoms may be directly related to the early availability of ascospores to act as a primary inoculum, as demonstrated for other ascomycete pathogens of plants, including *Z. tritici* (Morais et al., 2016), *Pyrenophora tritici-repentis* (Adee & Pfender, 1989) and *L. maculans* (Naseri et al., 2009). However, the observations reported here should be interpreted with caution, because flax seedlings emerged more than three weeks later in the 2016-17 season than in the 2014-15 and 2015-16 seasons, due to a lack of rain after sowing (**Figure Suppl. 2**). Moreover, weather conditions contrasted strongly between the three growing seasons at Thiverval-Grignon. In the period before flax crop

emergence in the 2016-17 season, July and August were particularly dry, with a total rainfall

of only 32 mm (Figure Suppl. 2). In 2014-15 and 2015-16, the period before autumn crop

emergence was less dry, with total amounts of rainfall for July and August rainfall of 151 mm

in 2014-2015 and 127 mm in 2015-2016. The growing season with the driest summer was the

least favorable for pseudothecium maturation, which was delayed by almost two months.

Discussion

This epidemiological study provides basic information about the interepidemic dynamics of the

flax pathogen M. linicola, the sexual stage of which was identified in all its forms -

pseudothecia, asci, and ascospores – directly on plant host tissues, for the first time in France

and in continental Europe. In England, Perryman et al. (2009) trapped airborne M. linicola

ascospores, but did not observe pseudothecia, raising doubts about the actual presence of the

sexual stage on flax residues, and concerning its quantitative significance in particular. Based

on our findings and other published evidence, we can now argue that the pathogen survives on

flax straw and that sexual reproduction plays a significant role in the epidemiology of the

disease. Our epidemiological data suggest that M. linicola ascospores produced on flax straw

are the major source of primary infection, at least in French conditions. The temporal dynamics

of pseudothecium maturity revealed that wind-dispersed ascospores could potentially initiate

pasmo epidemics as soon as flax seedlings emerge in the field, in October for winter flax and

February-March for spring flax.

Perryman et al. (2009) reported that epidemics began earlier in the growing season when there

was much more rainfall in the autumn (1997-98) than growing seasons with drier weather

(1998-99, 1999-2000). These findings are consistent with our own, which also suggest a

concordance between the earliness of the ascospore peaks (early September 2015 vs. late October 2016) and the earliness of pasmo symptoms (mid-November 2015 vs. mid-December 2016) in field conditions. The dynamics of the *M. linicola* sexual stage on flax residues during the interepidemic period and the pattern of change in pseudothecium maturity over time are consistent with the changes in the concentration of ascospores in the air measured by Perryman et al. (2009) in England and Scotland. Ascospore peaks have been recorded in September and October, the period when the proportion of mature pseudothecia is maximal. Our data, and the few published epidemiological findings available, suggest that ascospores probably play a major role in initiating epidemics, contrary to the prevailing view that pasmo is a seed-borne pathogen (Holmes, 1976).

Our data also suggest that precipitation and humidity are important for pseudothecium maturation, consistent with published findings for other ascomycete pathogens of plants. Temperature is known to affect pseudothecium maturation in ascomycetes, as established, for example, for *L. maculans* and *L. biglobosa* (Toscano-Underwood et al., 2003; Naseri et al., 2009), but this was not demonstrated here because the mean temperatures from early July to late October were similar in the three years considered (16.6°C in 2014-15, 16.5°C in 2015-16 and 17.3 in 2016-17).

The increase in the frequency of pasmo in France in the early 2010s, may, like the increase observed in the UK in the late 1990s (Perryman & Fitt, 2000), reflect an overall increase in the area under flax historically concentrated in the northern part of France, close to those of UK and Belgium (**Figure 5 and 6**). However, it may also reflect an increase in the cultivation of different types of flax crop, resulting in an almost permanent presence of host plants, acting as both a target (living and susceptible tissues) and a source (dead tissues, source of inoculum) of

the pathogen. Flax and linseed are sown in the autumn (for winter crops) or the late winter (for spring crops). Winter flax is rarely grown in France (< 1 %), with spring flax the most prevalent form cultivated (85%), followed by winter linseed (12%) and spring linseed (3%) (data CIPALIN; Labalette et al., 2011). In addition to the overall increase in the area under flax crops in France in the three last decades, the diversity of this crop and related practices (but also changes in these practices, e.g. management of straw during the intercropping period) may have account for the size of pasmo epidemics in growing seasons with favorable climatic conditions. First, a resurgence of pasmo in spring flax crops may have occurred over the last two decades in France due to the large amounts of ascospores produced on crop debris as a result of the increase in the area under winter linseed. Pasmo was of little importance on spring linseed before the 1990s in the UK, and Perryman et al. (2009) suggested that the disease would not have become a problem if winter linseed had not provided a source of inoculum. Indeed, straw is poorly degraded and difficult to plough under, since the fibers wrap themselves around disks, wheels and shovels. In the past, the only way to cope with linseed straw was to drop it in windrows after the combine and then burn it directly or harrow or rake it into piles and then burn it. In Canada straw choppers on new combines have been used to effectively chop and spread flax straw, if the straw was dry and relatively short or fiber content was relatively low (Flax Council of Canada, 2015). In UK and France, burning has been practiced for a long time. National regulations still allow to burn flax residues on the ground (e.g. Décret 2015-1769; Legifrance, 2015), but the practice tend to be reduced since the use of straw as biomass energy source is possible. All these changes in management of linseed straw may have played a role in the survival of *M. linicola*.

Second, the increase in pasmo levels in France coincided with a strong increase in the proportion of flax relative to linseed in 1995 (**Figure 5**). It is acknowledged that ploughing or burning crop debris which may carry *M. linicola* inoculum is important to control the disease

(Paul et al., 1991), but these practices make sense only for linseed because flax straw is left on the soil for retting (a process in which the action of microorganisms separates the fibers from the other parts of the plant) at the end of summer, sometimes until October. This process may, thus, make a significant contribution to the peak in *M. linicola* ascospore levels just before the emergence of the following crop. Finally, the 'brown bridge effect' during the interepidemic period (Kerdraon et al., 2019), boosting the overall amount of inoculum, may be particularly marked as flax and linseed are grown simultaneously in concentrated, close production areas (**Figure 6**), increasing the likelihood of detrimental interactions between sources of inoculum and target plant populations.

Third, resistance tests performed in controlled conditions within the framework of the 'SeptoLIN' project have highlighted significant differences in pasmo sensitivity between the main linseed (n = 7) and flax (n = 15) cultivars currently grown in France, the latter presenting higher levels of disease (Penaud et al., 2017). This significant difference (p < 0.05), assuming that it was stable over time, could also have contributed marginally to the increase in the frequency of pasmo.

This study illustrates how a particular part of the disease cycle, in this case, the sexual phase, can become crucial due to changes in agronomic practices and the processing of plant products. The epidemiological consequences of such changes to the production and processing system must be taken into account at larger spatiotemporal scales, to improve crop protection strategies. Several fungal diseases of agronomic importance caused by ascomycetes have a direct impact during the epidemic phase (yield reduction due to effects on plant growth) and an indirect impact during the interepidemic phase (increase in inoculum production and disease pressure at early stages in the development of the following crop). In the case of flax, there may be a double direct effect on both the period of cultivation and the interepidemic period, because sexual reproduction takes place on the valuable part of the plant – the stem, from which fibers

are extracted – during retting. Verticillium wilt, another important flax disease caused by the

soilborne fungus Verticillium dahliae, is known to damage flax fibers and to cause significant

yield losses. This fungal pathogen reaches the fiber during retting, leading to the embedding of

numerous microsclerotia within the bast fiber bundle of the stem, which becomes brittle and

fragile (Blum et al., 2018). Asexual infections of the stems with *M. linicola* do less damage to

the fibers than V. dahliae, and no specific impact of sexual reproduction (i.e. the formation of

pseudothecia within the fibers) has yet been established. However, the issue of possible damage

to the fibers due to pasmo is of interest. In particular, it would be useful to estimate the

biological, chemical and physical impacts of M. linicola on the retting process in the field, fiber

quality and the production of inoculum for the infection of flax plants in the following cropping

period.

We show here that the sexual stage of M. linicola is of great epidemiological importance in

pasmo. This importance may have increased over the last two decades. Our findings suggest

that the changes observed in flax production may have resulted in M. linicola injuring the plant

in two ways, depending on the production conditions (Savary et al., 2000; 2012) and

assessments of the yield losses caused by pasmo (Perryman & Fitt, 2000). Approaches taking

this damage into account pave the way for the maintenance of scutched flax fiber quality, and,

more generally, the sustainability of the French linen/linseed sector, through the control of

pasmo without the need for an increase in fungicide use.

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Figure captions

Figure 1 – A. Early septoria lesion on a flax leaf with pycnidia exuding Mycosphaerella linicola

cirrhi. B. Pycnidium, showing a cluster of several hundred pycnidiospores (×10). C. Flax

(Linum usitatissimum) straws collected from a field during retting. D, E. Pseudothecia (black

points) on flax stem residues. F. Mature pseudothecium (brown) and asci (×400). G. Ascus

containing 8 ascospores (blue).

Figure 2 – Sequence of experimental steps (Koch's postulates) demonstrating the relationship

between the presence of the sexual form of Mycosphaerella linicola on flax stem residues and

pasmo symptoms on leaves.

Figure 3 – Mycosphaerella linicola sexual fructifications. A. Pseudothecium; B Ascus; C.

Ascospores; D Pycnidiospores.

Figure 4 – Dynamics of the sexual stage of Mycosphaerella linicola on flax straw during the

2014-15, 2015-16 and 2016-17 growing seasons (from July to April), expressed as the change

in the percentage of mature pseudothecia over time (see Figure Suppl. 1).

Figure 5 – Change in the area under flax/linseed (flax for fiber production and linseed for oil

production) in France from 1970 to 2019 (data CIPALIN).

Figure 6 – A. Area (ha) under flax for fiber in the main producing French departments and in

the main neighbouring producing countries (UK, Belgium and the Netherlands) in 2019 (data

CIPALIN and FAOSTAT). B. Area (ha) under linseed in the main producing French

departments in 2019 (data CIPALIN).

Supplementary Figure 1 – Visual categories for recognition and classification of the

maturation of each pseudothecium of Mycosphaerella linicola based on the stage of

development of asci and ascospores (i.e. degree of maturity): A. Immature; B. Maturing; C.

Mature; D. Empty.

Supplementary Figure 2 – Mean daily air temperature (dotted line, °C) and daily rainfall

(histogram, mm) at Grignon from July to April in the 2014 (A), 2015 (B) and 2016 (C) growing

seasons, relative to the change in the percentage of mature Mycosphaerella linicola

pseudothecia on flax straws over time (black line). Cumulative rainfall in the summer, which is

thought to have a strong impact on the early dynamics of the sexual stage of M. linicola on flax

straw, is indicated.

Figure 1

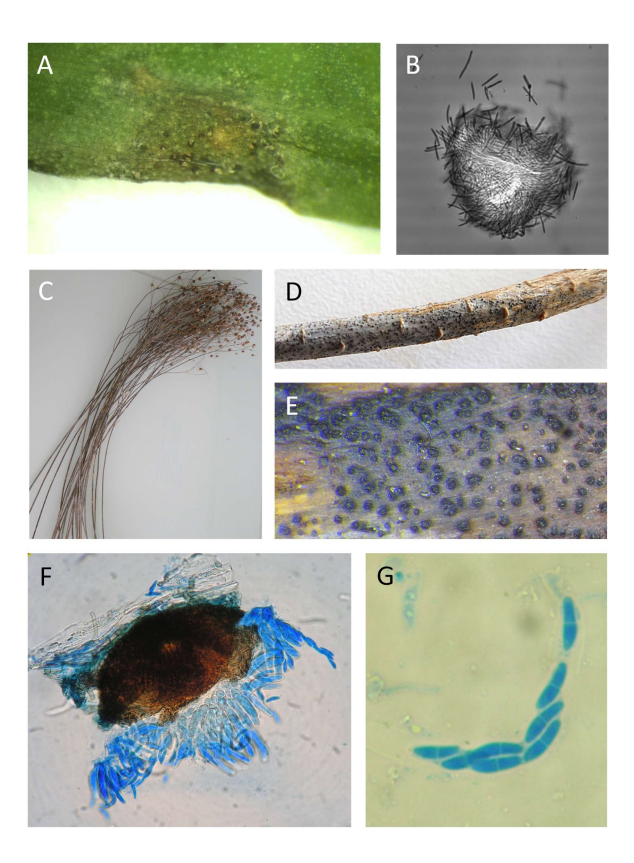


Figure 2

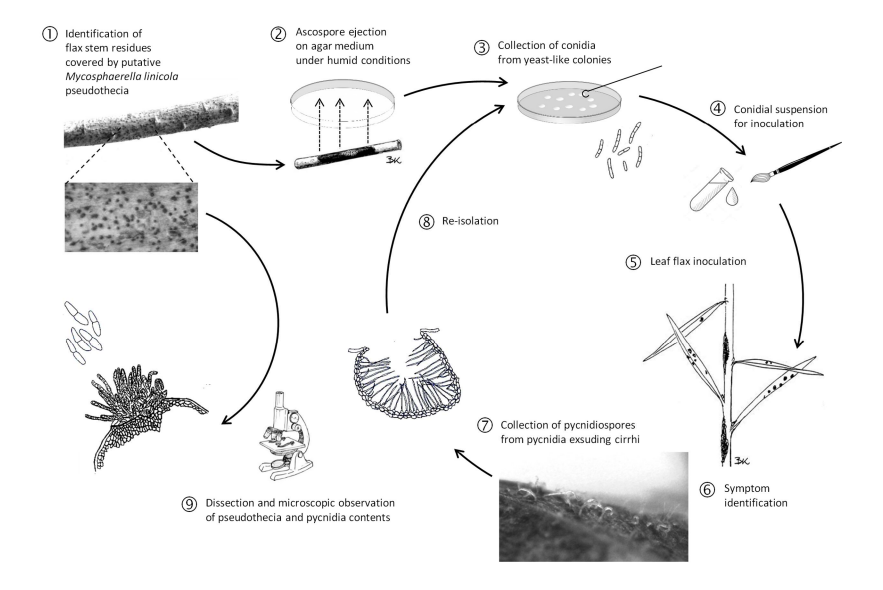


Figure 3

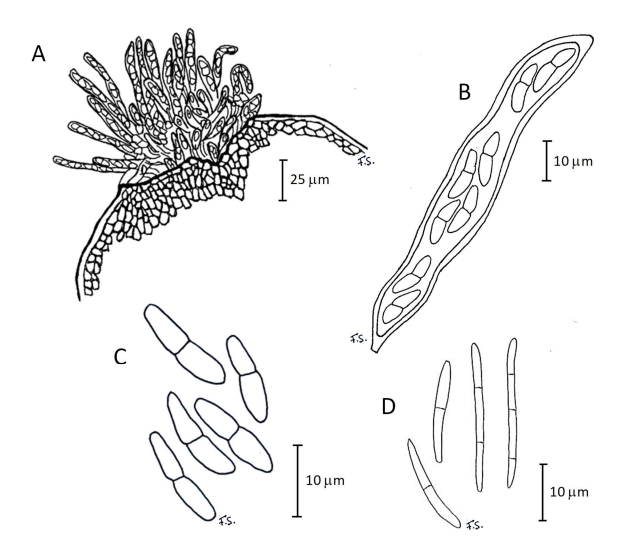


Figure 4

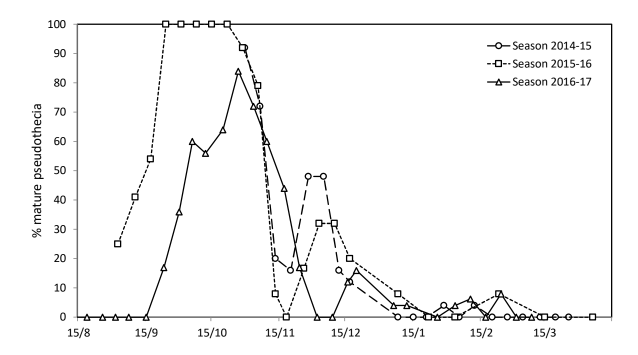


Figure 5

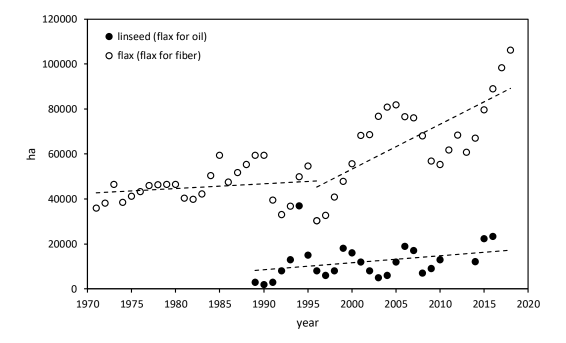
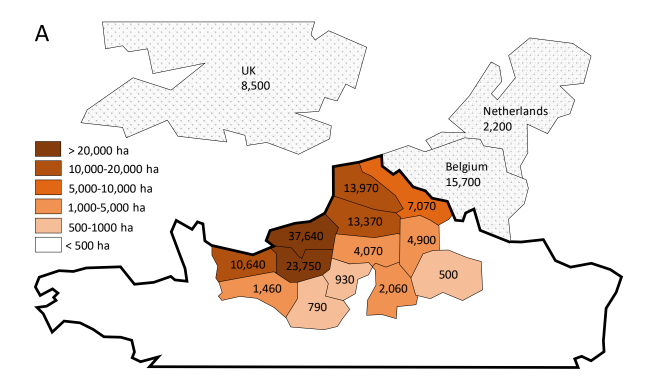


Figure 6



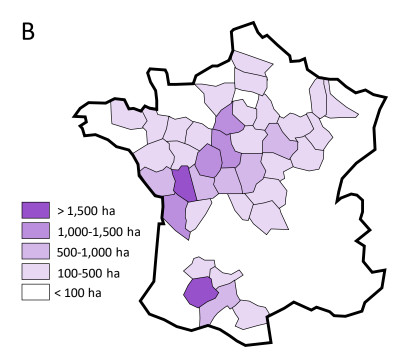


Figure Suppl. 1

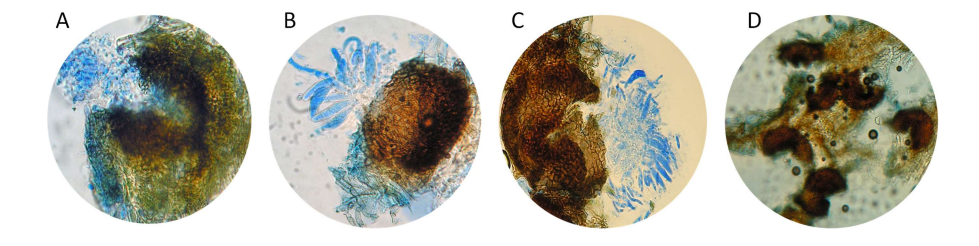


Figure Suppl. 2

