Fusarium head blight: epidemiological origin of the effects of cultural practices on head blight attacks and the production of mycotoxins by Fusarium in wheat grains

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Review — *Fusarium* head blight: epidemiological origin of the effects of cultural practices on head blight attacks and the production of mycotoxins by *Fusarium* in wheat grains

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Abstract

*Fusarium* head blight is an ancient disease and is very common throughout the world. In this article, we review current knowledge concerning the effects of cultural practices on the development of head blight and the production of toxins in the field. The qualitative effects of these practices on the severity of the disease and/or the production of toxins are in the process of being elucidated but, in many cases, detailed studies have not yet been carried out or conflicting results have been obtained. However, it should be noted that these effects have not yet been quantified. Three different cultural practices are today considered to be of prime importance for combating this disease and the production of mycotoxins: deep tillage, the choice of the preceding crop in the rotation and the choice of appropriate cultivar, as varietal effects exist.

Keywords: head blight, *Fusarium*, *Microdochium*, wheat, cropping systems, mycotoxins

Contents

Introduction ................................................................. 2
1- Epidemiological study of the disease ........................................ 4
2- Effects of various cultural practices on the disease ..................... 13
3- Effects of various cultural practices on the production of mycotoxins by *Fusarium* ............ 24
4- Relationship between the disease and toxin production .................... 28
Conclusion .................................................................................. 39
References ................................................................................... 44
Introduction

Head blight is a fungal disease affecting many small-grain plant species worldwide [1-10]. Affected wheat grains are small, light (the kernel is degraded), wrinkled and sometimes covered with a white or pink down. Rings or oval stains with brown edges and clear centres may be visible on the back of the grain [1,2,6,11-13] and on the external surface of the glumes (Figure 1). Head blight results from the development of a complex of two genera of pathogenic fungi: *Microdochium* and *Fusarium* [8,14-22]. *Microdochium* consists of a single species, *M. nivale*, made up of two subspecies: *M. nivale nivale* and *M. nivale majus*. *Fusarium* consists of five main species (*F. graminearum, F. culmorum, F. avenaceum, F. poae* and *F. triticum*), with several strains per species. The most common of these species are *F. graminearum* and *F. culmorum* [2,4,5,23-27], which are also the most pathogenic, reducing the size of the grains and the final yield [24,28].

The consequences of head blight contamination

This disease was first described at the end of the 19th Century in England [29] and has economic consequences. Yield losses (flower abortion, decrease in grain weight, highly damaged grains eliminated during threshing) of wheat may reach 15 stooks (180 sheaths) per hectare in cases of severe disease (according to the technical organisation dealing with cereals in France), or 50-60% [1,6,7,30] about once every three to five years [6,16]. In 2000, the year with the highest levels of disease in the last ten years, up to 100% of ears were affected on certain French plots [31]. There is a non-linear relationship between the percentage of ears attacked by *M. nivale* and thousand-grain weight [19,32]. Head blight also decreases the nutritive and technological quality of the grains (degradation of proteins, baking quality, nutritive value etc.).

Since the 1990's, interest in food health and safety has increased. For cereal products, one of the most important elements of alimentary risk is the possibility of mycotoxin accumulation on or in the grains [33-35]. Mycotoxins are the products of fungal secondary metabolism [34,36], which occurs when growth stops altogether or slows markedly [37]. These toxins result from adaptation of the growth of the fungus to stressful situations [37]. Head blight is the principal means by which mycotoxins develop in the grains before harvest. The mycotoxins produced in crops suffering from head blight are produced by fungi of the genus *Fusarium*. Fungi of the genus *Microdochium* are thought not to produce toxins [8]. Levels of
the mycotoxins produced by \textit{Fusarium} cannot be reduced after harvest by means of classical transformation procedures [38]. These toxins are chemically and thermally stable [39,40]. They may therefore be found in the raw material (grains) and in transformed products (flour, bread) or in products from animals fed with contaminated grain (meat, milk, eggs) [35].

Mycotoxins cause two types of problem. They may generate technological problems, such as negative effects on malting for beer (inhibition of enzyme synthesis) or on fermentation (inhibition of yeast growth), or even on the production of high-quality bread [38]. Indeed, \textit{F. graminearum} modifies a protease that acts on gluten. Damage to this enzyme prevents the dough from retaining the gas produced during fermentation, resulting in heavier, less voluminous bread. Furthermore, \textit{F. graminearum} also destroys starch grains, protein stores and cell walls [6,41,42].

Mycotoxins may also cause poisoning incidents, the principal symptoms of which are nausea, lethargy, fatty and cancerous infiltrations and possibly even death [38,43]. Six main families of mycotoxins have been studied to date: aflatoxins (B1, B2 and M1), ochratoxin A, trichothecenes (T2 toxin, deoxynivalenol etc.), zearalenone and fumonisins. \textit{Fusarium} produces toxins of the last three groups, which may attack the liver, the kidneys, the nervous system, the circulatory system, the endocrine system, the skin, the digestive tract and the blood [44]. These toxins are thought to be highly carcinogenic, although this assertion remains unproven in most cases. However, T2 toxin has been shown to be carcinogenic [44].

The absorption of trichothecenes has resulted in the poisoning of animals and humans in the United States, Canada, the Soviet Union (as it was at the time) and Japan [39]. The massive absorption of T2 toxin was responsible for toxic alimentary aleukia in several regions of the USSR between 1942 and 1947 [34,39]. Pigs are the animals most reactive to deoxynivalenol, whereas poultry seem to be particularly resistant. Effects on humans have been observed after the absorption of 300 to 8000 µg/kg deoxynivalenol in India [8]. This led the public authorities to establish norms fixing maximum deoxynivalenol levels in food destined for human consumption. The European Commission has proposed setting limits of 750 µg/kg (750 ppb) in cereals and 500 µg/kg in cereal-based products such as flour. A major problem in trichothecene toxicology is the current lack of knowledge concerning the risk of food contamination by several of these toxins at the same time, as \textit{Fusarium} can produce a multitude of toxins simultaneously [34].
**Factors triggering attacks**

The climate is known to have an effect, and is said to be the principal factor affecting the development of head blight on oat in Norway [45]: humidity determines the severity and intensity of the disease whereas precipitation and total radiation determine inoculum levels (number of spores per ear) [46]. The climate plays a role at all stages of development of the fungus [47-49]. The climate during winter affects the survival of the fungus on crop residues: during mild periods of the winter, the fungus is capable of sporulating on the cereal stubble debris [48]. In spring, the climate determines the type and quality of spores produced, together with the date of their dispersal and thus the intensity of the disease. Indeed, coincidence in the dates of spore dispersal and crop flowering, marking the start of the critical phase of infection for plants [2,6,16,50], is of prime importance in determining the intensity of the attack. In summer, together with the virulence characteristics of the fungal strain and the intrinsic resistance of the plant, climate controls competition between pathogen species, thereby controlling the rate of mycelium growth in the plant [47]. The climate may also affect the production of fungal toxins in the field [2,51-54]. The effects of climate are such that the levels of deoxynivalenol recorded on diseased crops in the field and in the glasshouse are not correlated [49]. The effects of climate are of course uncontrollable, but also difficult to predict due to their complexity.

However, in any given year, two neighbouring plots may display different levels of infection [19,55]. It must therefore be possible, to some extent, to control the production of mycotoxins by modifying the cropping system.

The literature review below aims to identify and to elucidate the relationships linking cropping systems to grain contamination by *Fusarium, Microdochium* or to mycotoxin production by *Fusarium*.

**1- Epidemiological study of the disease**

**1-1. Impact of climate on the species complex**

The species of fungi making up the pathogenic complex responsible for head blight may differ from year to year and from one region to another, particularly as a function of climate [6,11,56] and wheat variety [56,57]. Indeed, climate partly controls competition between the
various species. Thus, at flowering, the development of *Microdochium* is favoured by lower temperatures [6,58] and rainfall [59], whereas that of *Fusarium* is favoured by high temperatures [58] and storms [59]. *F. graminearum* is generally the predominant *Fusarium* species in warmer regions, whereas the predominant species in cooler regions are *F. culmorum* [6,16,28,60], *F. avenaceum* [6] and *M. nivale* [6,16]. Furthermore, in spring (before ear emergence), drought favours the development of symptoms of *F. graminearum* and *F. culmorum* infection at the base of the plant, at the expense of other species of *Fusarium* [59].

1-2. Sources of inoculum

Several reservoirs of the parasite complex responsible for head blight are known. The primary reservoir of inoculum is debris from the previous crop [2,61]. All species of *Fusarium* and *Microdochium* can survive as saprophytes [6]. However, the pathogens survive longer on residues that do not degrade easily, such as stem nodes [2].

Other sources of inoculum include numerous plant hosts. These may be cultivated plants and/or weeds, such as grasses and evergreen weeds [6]. However, the roles of weeds, inoculum source (site at which the fungi develop) and supports for survival have not yet been determined [62]. Seeds are the other major source of inoculum, making it possible for the disease to begin in autumn in the case of grains contaminated with mycelium [14,16]. The further into the grain the mycelium has penetrated, the lower is the chance of the grain germinating. The soil may also be contaminated [1,2]. Soil-borne infections take hold less rapidly than seed-borne infections, resulting attacks affecting essentially the collar and the upper parts of the roots [2,14]. Soil humidity, particularly during winter, decreases the pathogen survival rate [2].

The flag leaf is the principal site of spore production and source of ear infection in the plant [32].

1-3. Production and maturation of inoculum

During the winter, the fungi survive as chlamydospores, mycelium or propagules [2,6]. Sporulation, which involves the production of various types of spore, occurs during mild periods [48]. Four types of asexual inoculum may be produced, depending on the species concerned: macroconidia, microconidia, chlamydospores and hyphal fragments, the size and
form of which depend on the species of pathogen [63]. The macroconidia of *F. graminearum*
may contain one to seven septa — most frequently three to seven — and measure 20 to 105
µm in length (mostly 35-62 µm), and 2 to 56 µm in width [2]. This type of inoculum may be
available during the entire crop cycle [16]. Sexual spores, known as ascospores, may also be
produced by reproductive organs called perithecia. Only three of the species most frequently
responsible for head blight appear to be capable of producing ascospores in the natural state:
*M. nivale*, *F. graminearum* and *F. avenaceum* [6,16,64]. These three species produce
ascospores that are similar in size and form. They generally have three septa and measure
about 17.5 – 26 µm x 3.5 – 5 µm [2]. They take a long time to mature and therefore
contamination cannot occur before ear emergence [16]. Caron reported that *M. nivale*
produces ascospores from May onwards in France and that *F. graminearum* produces its
ascospores later in the year.

In the laboratory, the optimal environmental conditions for production of the principal forms
of inoculum — macroconidia and ascospores — depend on the species and the environment.
These results probably cannot be directly transposed to natural conditions. Table 1 sums up
the principal results reported in the literature.

The maturation of the inoculum has been studied in less detail and depends on interactions
between environmental factors, both in the laboratory and in the field. The maturation of the
inoculum is thought to be hindered by drought and cold in autumn and winter [65]. In the
laboratory, perithecia have been reported to mature in six to nine days [66], or nine to ten
days [2] after their initiation in ideal conditions. In contrast, perithecium maturation takes two
to three weeks in the field whereas conidiospores can be produced in large quantities within a
few hours [59]. According to another study, macroconidia reach maturity in three weeks in
the field [67]. The rate of maturation of perithecia depends on light [66,68].

1-4. Dispersal of the inoculum

The inoculum is dispersed by various animal vectors. Mites (*Siteroptes graminum*) transport
*F. poae* spores [6]. Insects are parasitised by certain species of *Fusarium*, *F. episphaeria* (SN.
and H) in particular [63]. *F. avenaceum*, *F. culmorum* and *F. poae* have been isolated on
various insects including *Musca domestica* (housefly), *Hypera punctata* (clover leaf weevil)
and *Melanoplus bivittatus* [69].
The possibility of inoculum dispersal by the systemic route has long been debated; after cutting the peduncle of a wheat ear into segments, Atanasoff (1920)[1] observed *F. graminearum* in the segments close to the ear but not in those close to the flag leaf. Following inoculation of the base of the wheat stem, only 3% of plants display colonisation beyond the second node and no fungus is detected beyond the fifth node [70]. In another study [71], the tops of plants produced from seeds inoculated with *M. nivale* presented similar numbers of perithecia to those of plants grown from healthy seeds, even though the plants grown from contaminated seeds had more perithecia at the base of the stem. This finding confirms the lack of relationship between head blight and foot rot due to *Fusarium* [6,16]. To date, these two infections have been considered to be essentially independent [1,16]. However, it should be noted that the presence of the parasite at the base of the stem (below the second node) may disturb the water supply to the ear sufficiently to cause shrivelling at the slightest increase in temperature [14]. Shrivelling of part of the ear is one of the symptoms of head blight. A confusion of symptoms is possible: in some cases, shrivelling of the ears may be due to the presence of the parasite at the base of the stem rather than on the ears. In addition, following the artificial inoculation of spring wheat seeds, sown in pots, with *F. culmorum*, the pathogen is isolated from all nodes and from the ears [72]. Similarly, Snijders (1990a)[73] observed the pathogen in stem tissue 70 cm above soil level after inoculating plants at soil level. Many cases of infection with various species of *Fusarium* have been observed after the sterilisation of weeds [62] and wheat [74] with sodium hypochlorite, which suggests that these infestations are endophytic. Evidence that this is indeed the case is provided by the observation that the fungal mycelium is capable of infecting both the parenchyma and the vascular tissue, in which it is able to travel more rapidly [50]. Indeed, after the injection of spores into the rachis, these authors showed that the pathogen was able to migrate within the plant and that it propagated more rapidly longitudinally than transversely. However, it is possible that this route of contamination is only possible in a few species of plant, for a few species of fungus or for a few plant-fungus interactions.

Dispersal by leaf-to-leaf contact also seems likely as the pathogens (*Microdochium* and *Fusarium*) are found on the leaves [15]. Atanasoff (1920)[1] also observed infection by contact. This mode of dispersal should be considered to be a specific case of aerial contamination.
Aerial contamination by ascospores and conidia is thought to be the principal source of ear contamination [1]. This means of contamination has been studied by many groups and involves two possible modes of dispersal: splashing and wind [2,6,7,16,64]. The relative importance of these two modes of dispersal depends on the climate and the species making up the pathogenic complex in the year and region studied and the capacity of these species to produce ascospores.

Splashing transports spores, macroconidia in particular, that are too heavy to be transported by wind [35]. The density of *F. graminearum* macroconidia in the air above the canopy has been estimated to be only one twentieth that of ascospores from the same species [75].

Splashing is the only means of dispersal of *M. nivale* conidia [71]. Millar and Colhoun were able to trap conidia only during simulated rain conditions, whereas Fernando *et al.* (2000)[75] observed a peak in the release of macroconidia of *F. graminearum* in the air one to two days after rainfall following a long period of drought. These two studies demonstrate the major role played by rainfall in the dispersal of spores. In the laboratory, a conidium of *F. graminearum*, produced in a sporodochium (the asexual fruiting body bearing the conidiophores) from wheat straw, receiving a single drop of water 5 mm in diameter falling from a height of 6 m, was displaced by up to 45 cm in the vertical plane and 90 cm in the horizontal plane [17]. In the same conditions, a conidium of *F. culmorum* was displaced 60 cm vertically and at least 1 m horizontally [17,76]. The spores of *F. poae* may reach a height of 58 cm and may travel 70 cm horizontally in rebounding raindrops [76]. Splashing alone is therefore sufficient to transfer a conidium from crop residues or the stem base to the ear (in one or several rebounds, relayed by the leaves), assuming there is no obstacle. In the canopy, the leaves of the neighbouring plants form an obstacle. Under simulated rainfall, the wheat canopy reduces the dispersal of *Septoria tritici* spores by 33%, to 15 cm, in the horizontal plane from the source of infection, and by 63% in the vertical plane [77]. A leaf positioned low in the canopy is principally infected at its base whereas leaves higher up in the canopy are principally affected at their tips, from which the spore can rebound, resulting in their transfer to another leaf [76].

Splashing is considered to be the most likely means of dispersal because *F. avenaceum*, *F. culmorum* and *M. nivale* have been observed on the flag leaf [15]. Rainfall plays an important role in the development of the disease. In years with major epidemics of head blight or rotting of the ear due to *F. graminearum*, precipitation levels are generally high [2,7,15,78]. Similarly, in spray irrigation trials, up to 89% of ears may be infected, versus 0% in non-
irrigated controls [6]. Furthermore, humidity and rainfall in spring favour the formation of perithecia [59].

In natural conditions, perithecia are formed in only the three species that generate ascospores: *F. graminearum*, *F. avenaceum* and *M. nivale* [6,16,64]. Ascospores are a form of inoculum that can be transported by the wind [6,16]. The wind has long been considered the principal vector for spore dispersal, and damage tends to be greatest in the direction of the prevailing wind [1]. Following the artificial inoculation of a zone of field with maize grains infected with *F. graminearum*, *F. graminearum* spores travel less than 5 m [75]. However, Stack (1997)[79] indicated that the dispersal distance of *F. graminearum* spores is proportional to the size of the area inoculated. According to this model, head blight levels halve as the distance to be travelled by the inoculum doubles. The maximum density of ascospores observed is 1500 spores/m², at 1.5 m from the source of inoculum [75]. The dispersal of *F. graminearum* ascospores in the field occurs at temperatures of 13 to 22 °C, with a relative humidity of 95 to 100% [2]. Ascospore release is maximal at a relative humidity of 100%, but also occurs at lower levels of relative humidity [66].

Despite the importance of humidity, studies on *F. monoliforme* [67,80] and *F. graminearum* [75] have shown that the release of ascospores is not directly linked to rainfall. Indeed, ascospore release peaks one to four days after rainfall [67,75] of at least 5 mm or a relative humidity exceeding 80% [67]. Paulitz found that a shower of rain in the evening did not inhibit spore release but that heavier rain (at least 5 mm), a relative humidity exceeding 80% continuously throughout the day or the alternation of rain and high humidity inhibited the release of ascospores during the night. This result contrasts with another study showing the release of ascospores during the night following a day on which 8 mm of rain fell [81]. The correlation between spore release and humidity suggests that the maturation of perithecia requires a certain level of humidity. This would account for the results of Fernando *et al.* (2000)[75], who observed four periods of spore release over a period of 20 to 30 days.

The process of ascospore release may be described as follows [71]: after a period of drought (air with low water vapour content), free water (rain or heavy dew) triggers the release of ascospores. Paulitz (1996)[67] speculated that the increase in relative humidity during the evening after a dry day might increase the turgor pressure of the asci, which contain a vacuole at high osmotic pressure [82]. This high osmotic pressure is generated by the accumulation of
mannitol and the flux of potassium and calcium ions [66]. Calcium ions are also involved in
signalling for ascospore discharge [66]. This increase in pressure triggers the release of
ascospores, which are expelled from the perithecia in a gelatinous substance, which then dries
out, releasing the ascospores into dry air. This would account for the finding that a trap placed
above the canopy captures several *M. nivale* ascospores in dry conditions and that the
concentration of these spores increases by a factor of 20 in the 10 minutes following the
application of a fine simulated dew [71]. A similar phenomenon has also been observed for
the ascospores of *F. graminearum* [6]. Similarly, in control conditions, the maximal release of
*F. graminearum* ascospores is regulated by dehydration of the perithecia in non-saturated
humidity conditions [83]. Furthermore, the time at which the spores are released is strongly
correlated with the increase in humidity following the decrease in temperature that occurs at
the end of the afternoon [67]. Paulitz observed that ascospore release began before the leaves
became humid at the base of the canopy (which occurred at 22-24 h), towards 17 or 18 h for
temperatures varying between 11 and 30 °C and values of relative humidity of between 60
and 95%. These results have been confirmed by the observation, in control conditions, that
the maximal release of *F. graminearum* spores occurs at temperatures between 11 and 23 °C,
with 16 °C the optimal temperature [83].

After temperature and humidity, the next most important environmental factor affecting
ascospore release is light. Light is required for the production and maturation of perithecia
[68] and ascospores [83]. In contrast, the process of ascospore release does not directly
require light, as it has been observed during the night: 4 to 5 h before sunrise [67]. This result
has been corroborated by studies in control conditions, which also showed that light had no
effect on ascospore release [66,68].

Finally, the periodicity of aerial dispersion has been studied. The release of conidia is not
periodic, whereas that of ascospores is, particularly in *F. graminearum* [75]. Indeed, Fernando
*et al.* detected no ascospores in the air between 12 and 16 h, whereas ascospore density was
maximal between 20 h and 08 h (1.5 times higher than at other periods of the day). Other
authors have reported that ascospore density peaks between 21 h and 08 h [84]. Similarly,
Paulitz reported that spore release often occurs between 18 h and early morning (04 h – 08 h),
mostly before midnight, with a peak at around 23 h [67]. For *F. graminearum*, the maximum
ascospore concentration observed is of the order of 4333 ascospores/mm³ in one hour. Paulitz
also reported spores of other species of *Fusarium* to be continuously present in the air, but
with densities varying according to the period of the day. Paulitz identified four distinct periods in the day, and found that spore density was low in the morning. For *F. monoliforme*, a large number of ascospores was found to be released between 17 h and 08 h, with a peak towards 02 h [80].

Although dispersal over large distances seems to be possible, local aerial contamination appears to predominate, in that two neighbouring plots of land planted with the same variety may display very different levels of contamination [19,55].

1-5. **Infection and colonisation of the ear**

In wheat, the most critical period corresponds to the moment at which the degree of resistance to primary infection and to the propagation of fungal hyphae in the plant is lowest. This period has been defined differently by different authors, but the first half of the grain-filling period is the most critical. Several authors have identified ear emergence [64,85] or the mid-grain-filling stage [47] as the most critical, but most authors consider anthesis to be the most susceptible to attack in wheat [12,16,64,86,87], with susceptibility decreasing strongly after the start of the dough stage [12,16,47,64,86,87]. Elimination of the male organs from wheat ears decreases the frequency of infection by *F. graminearum* [47,86]. Similarly, sterile wheat lines are less susceptible to head blight than fertile lines [88]. This, together with extensive colonisation of the anthers by this fungus [86] indicates that the growth of *F. graminearum* is stimulated in these structures, suggesting that entry into the anthers during anthesis has major consequences for the grain (degradation). Two substances — choline chloride and betaine hydrochloride — are much more concentrated in the anthers than elsewhere [89]. These substances favour the extension of conidial hyphae, but not the germination of spores of *F. avenaceum*, *F. culmorum* and *F. graminacearum* [89]. Once the spores have germinated, the propagation of the hyphae is therefore more strongly favoured in the flower parts than in the other organs. However, the difference in susceptibility of different stages is probably due to the fact that the critical receptivity peak observed (degree of resistance to primary infection and propagation of fungal hyphae) is dependent on many factors. These factors include the wheat cultivar [12,47,50,90], air temperature and humidity [12,47] at the critical stage for the plant and, before this stage, development of the inoculum, and the stage of maturity of the plant at the time of hyphal penetration [12]. The testa (seed coat) is more resistant to penetration when the grains are mature [12]. However, this finding remains open to debate because a recent study [91] showed that the date of inoculation has no effect on the final level
of *F. graminearum* colonisation. In maize, susceptibility to the disease is maximal at the start of silk development and decreases throughout the grain-filling period [92]. In *Arabidopsis thaliana*, tobacco, tomato and soybean, the floral tissues are also the most heavily infected [93].

The infection process is very similar in susceptible and resistant varieties [94]. The pathogen first penetrates host tissues 36 to 48 h after inoculation [94]. The first organs affected are the lemmae and the tip of the ovary [94,95], the anthers [12] and/or the spikelets, glumes and the rachis [50]. It remains unclear which of these organs is most frequently the initial zone infected, but this probably depends on the date of infection and possibly also on the infecting species. The penetration of the fungus into the ear is favoured by relatively low temperatures and high humidity [64]. The presence of and colonisation by a large number of hyphae are required for infection [50]. The hyphae of *F. graminearum* and/or *F. culmorum* invade the host tissue directly [94] or via the stomata [50]. They then propagate into the ear, passing through and around the cells in their path [12,94-96], degrading the cells that they contaminate [50,94,96]. They move principally towards the rachis [94,95] or towards the young grains, which they invade via the parenchyma of the pericarp close to the embryo [50]. A short time after flowering, the parenchyma of the pericarp begins to break down, the nuclei and cytoplasm of the cells disappear and the walls of the cell break [12]. *F. graminearum* is then able to enter this tissue and propagate throughout the grain [12]. Indeed, *F. graminearum* hyphae penetrate the thinner cell walls of the parenchyma tissue more easily than the thicker cell walls of other more specialised tissues [12]. Birds also facilitate infection by creating lesions on the grains that favour the penetration of fungi [2].

Infection of the ears by macroconidia of *F. graminearum* [47], *F. avenaceum*, *F. culmorum*, *F. poae* and *M. nivale* [6] is optimal at 100% relative humidity and 25 °C, and takes place over 24 to 60 hours. Infection may occur at temperatures of 20 to 30 °C, but is negligible at temperatures below 15 °C [47]. Below 18 °C, the conidia of *M. nivale* are more competitive than those of *Fusarium*, resulting in higher levels of *M. nivale* contamination. Furthermore, a period of four to eight days of low humidity reduces the incidence of infection, but does not entirely eliminate it [47]. The germination of macroconidia, ascospores and chlamydospores is maximal between 0 and –20 bars and is inhibited between –60 and –80 bars [97]. Ascospore germination is inhibited beyond a threshold of –30 bars (-3 MPa) of water.
potential after eight hours of drought [67]. Free water or a low water potential also favours infection, reducing the length of the incubation period [67].

1-6. Incubation and sporulation
Perithecia and conidia develop on the surface of spikelets and of the rachis in humid climatic conditions [2]. The duration of the incubation period required decreases with increasing humidity [16]. In conditions of saturating humidity, the duration of incubation required for the appearance of macroconidia of *F. culmorum* and *F. graminearum* on the ear was 12 days at 14 °C, less than five days at 20 °C and less than three days at between 25 and 30 °C [2,16]. More spores are formed after a long period of high humidity. This may then result in the infection of later crops, such as maize.

2- Effects of various cultural practices on the disease
Various studies have identified different elements concerning the effects of agricultural practices (crop rotation, crop management) on head blight attacks.

2-1. Effects of the crop succession history of the field
The effects of crop rotation have been studied in detail. They depend on the preceding crop, whether that crop is a potential host for the pathogens responsible for head blight, and the frequency of the crop concerned in the rotation. The shorter the rotation, the higher the frequency of head blight. Thus, head blight is most frequent when the susceptible crop occurs frequently within the rotation [7]. The density of crop residues left in the field [19], their nutritional value [19] and pathogen competition may also modify the effect of crop rotation [19,62].

Head blight contamination is more severe if the preceding crop is maize, durum wheat or oats, rather than wheat or barley [14,19,48,64,98-101], and even less contamination is observed following other crops [16,48]. For example, the frequency (%necrotic ears) and the severity (%necrotic spikelets) of the disease on wheat are lower following soybean than following another wheat crop, or worse still, maize [19]. This may be accounted for in two ways. Firstly, soybean crops leave fewer residues than wheat crops, which in turn leave fewer
residues than maize crops [7,19,100]. Secondly, the principal *Fusarium* species infecting soybean is *F. sporotrichioides*, whereas wheat and maize are more frequently affected by *F. graminearum* [19], like sorghum [59]. Thus, in addition to the density of residues left by the preceding crop, the nature of the preceding crop is important in determining the pathogen species likely to infect the next crop in the rotation. The preceding crop affects the composition of the pathogen complex throughout the following year. Thus, a preceding crop of potato will allow the development of only *F. culmorum* and *F. sambucinum*, whereas a preceding wheat crop will allow the development of all *Fusarium* species [62]. It should also be noted that the highest levels of *F. graminearum* contamination are recorded on grains harvested from wheat crops following maize in the rotation, whereas *F. avenaceum* and *F. poae* are the most common species found in grain samples harvested from wheat crops following a crop other than maize [60]. It is therefore advisable to introduce non-host plants into crop rotations, to limit the disease [7]. Indeed, the use of oats as the preceding crop results in a doubling of head blight inoculum in the soil [64,98] even though this crop is unaffected by the disease [14] whereas a preceding crop of sugarbeet, which is hardly affected by *Fusarium*, halves the incidence of the disease in the subsequent wheat crop [14,64,98]. Similarly, flax can be used as a clean-up crop [16], as can alfalfa, after which no disease is observed [102]. However, the quantity of crop residues and the development of pathogens on the preceding crop are not the only explanations of the effect of preceding crop. Wheat crops present different levels of infection depending on whether they follow wheat or durum wheat in the rotation, even though these two preceding crops produce similar amounts of residues and are equally permissive for *Fusarium* development.

Among the other elements possibly involved, the amount of nitrogen in crop residues may affect the possible duration of colonisation by *Fusarium*. Indeed, the high nitrogen content of maize residues may result in a longer period of colonisation of those residues by the fungi, strengthening these fungal populations [2] and favouring disease.

It is also likely that proteolytic soil bacteria, favoured by the low carbon to nitrogen ratio under alfalfa, compete with *Fusarium* species, limiting their development and thus their ability to attack the subsequent crop [62,102]. It should also be noted that, at the end of the cropping cycle, buried cereal stubble provides an ideal substrate for saprophytes, increasing the carbon to nitrogen ratio, possibly resulting in the reappearance of the disease [98].
2-2. **Effects of soil tillage**

The effects of soil tillage are also well known, at least from a qualitative point of view. As for yellow and brown rusts [103], limited soil tillage increases the frequency of head blight [7, 19, 100], whereas deep tillage (ploughing) decreases it [7, 19, 64, 90, 101]. Similarly, limited tillage systems increase the number of *Fusarium* propagules in the soil [14] whereas ploughing decreases inoculum levels [98].

This effect of ploughing may be accounted for in several ways. Ploughing has several direct and indirect effects on the structure [104, 105] and microclimate of the soil [105, 106] and therefore on the development of fungi [105], notably those of the genus *Fusarium*. Limited tillage and direct drilling systems make it harder to deliver the seed to the required position during sowing, decrease soil porosity, increase structural stability and litter on the surface. The litter layer protects the soil from rain (battering and erosion), and increases surface humidity (by limiting evaporation), soil temperature and inertia. This layer breaks down into organic matter, enriching the surface soil in carbon and organic forms of nitrogen, and modifying its chemical and biological characteristics. Stores of inorganic carbon increase, the supply of soil mineral nitrogen decreases very slowly and the mineralisation process is delayed. The microbial biomass (bacteria and fungi) increases at the surface and its turnover rate increases in the first 10 cm of soil. Humus is thus less well degraded in this zone, which rapidly becomes more acidic, favouring the development of fungi over that of bacteria. Thus, about 90% of the *Fusarium roseum* population is located in the first 10 cm of soil [98]. Furthermore, the development of this fungus depends on soil aeration [14]. Although this pathogen can survive for four years [16] at a depth of 20 to 25 cm, it is only active and able to develop on plant debris in the first 5 cm of soil [14]. Beyond a depth of 15 cm, the incidence of *Fusarium oxysporum* attacks on carnation decreases as the depth of propagule burial increases [107]. However, this result cannot be readily transposed to other species because *Microdochium nivale* and *Fusarium avenaceum* do not produce chlamydospores [108] and the number of propagules in the soil does not necessarily affect *Fusarium* foot rot levels due to *Fusarium culmorum* [109]. Thus, for certain species, propagules may simply ensure the survival of the fungus rather than serving as an inoculum [98, 109].

The development of the *Fusarium roseum* population also depends on the quantity of substrate (crop residues) available [14]. Thus, a decrease in the density of residues on the surface of the soil (after ploughing for example) helps to decrease the production of inoculum...
and the quantity of spores available for dispersal [109]. This hypothesis seems to be validated by the observation that there are fewer *Fusarium culmorum* propagules if the stubble from the preceding crop is burnt than if it is buried [109]. However, an opposite effect on the disease is observed. Thus, in years in which there are high levels of *Fusarium* infection at the base of the stem, disease levels have been found to be higher if the stubble from the previous crop was burnt than if it was incorporated into the soil [109]. A similar result was previously reported for eyespot [111]. Similarly, the supplementary incorporation of infected stubble not originating from the preceding crop decreases the incidence of *Fusarium* attacks resulting in wheat foot rot to levels below those observed in soils in which the density of residues has not been modified [112]. However, these results may be explained if stubble limits conidium dispersal by splashing, favours growth of the canopy [111], or modifies the infection process. In Yarham's study [111], the addition of stubble did not affect leaf area index, photosynthetic potential or the number of leaves. It also resulted in no decrease in the density of wheat or weeds and had no effect on the size of the plants. All these factors might have facilitated spore dispersal. In the infectious process, four aspects may be affected by the presence or absence of stubble. Firstly, the resistance of the plants may be affected by increases in the amount of silica (SiO$_2$) in the leaves of the wheat plants following the incorporation of additional stubble [112]. The hypothesis that silicon plays a protective role arises from work [103] indicating that this element strengthens cell walls and, particularly, the outer membrane of epidermal cells, increasing resistance to the penetration of pathogen germination tubes in leaves containing large amounts of silicates (shown experimentally for the penetration of *Erysiphe graminis*, which causes powdery mildew on barley) [113]. Similarly, adding silicon decreases *Fusarium* infections and powdery mildew in cucumber [114]. All changes in the structure and microclimate of the soil, the date, method and depth of stubble incorporation also influence the extent and dynamics of crop residue decomposition [105], which may in turn affect head blight levels. Indeed, in optimal conditions of microorganism colonisation and water and nutrient exchange, the area of contact between the soil and crop residues (affected by the size of the residues and soil porosity) is maximal, accelerating residue decomposition [105]. Consequently, despite providing a larger source of inoculum, stubble incorporation increases the rate of degradation of crop residues, decreasing the duration of colonisation of these residues by pathogens such as *Pseudocercosporella herpotrichoides* [110]. These pathogens are thus less well developed, resulting in lower attack rates, which in turn results in slower spore maturation following changes in humidity, light levels or temperature, or lower levels of spore production [111]. Indeed, it is possible that the
decomposition of stubble results in the production of substances toxic to fungi or that favour the microorganisms engaged in stubble decomposition, thereby creating competition between microorganisms [111], limiting the production of spores. Finally, a study of *Fusarium* foot rot showed that the distribution of crop residues is the primary factor influencing the site of penetration of the fungi [115].

2-3. Effects of irrigation

The irrigation of a field influences its microclimate and may encourage the development of the pathogen. Regardless of whether the climate is favourable for the disease in a given year, irrigation increases the frequency (% necrotic ears) and severity (% necrotic spikelets) of the disease over that in non-irrigated plots [19,90].

Uncertainties concerning the survival of the fungus in the soil and its capacity to sporulate in the soil as a function of soil aeration, porosity and light levels, together with the role played by the rate of residue degradation and that of the compounds generated by this process, and their mineral composition (nitrogen, silicon) limit our ability to make quantitative predictions of the effects of the preceding crop, soil tillage or irrigation.

2-4. Mineral nutrition

The effects of mineral nutrition on head blight attacks are unclear. As nitrogen applications favour the development of the plant, making the canopy more humid, it has often been suggested that such applications are likely to favour the appearance of diseases [103,116]. This has been demonstrated for bunt [103] and blotch or *Septoria* disease [117]. In fact, the severity of *Septoria* attacks on wheat is increased by 11%, which corresponds to a yield loss of 8-9%, following treatment with 100 kg of nitrogen per hectare [117]. Nitrogen application significantly increases the incidence of *Fusarium* infection in grains of wheat, barley and triticale [118,119] and the predisposition of wheat to attacks by *F. avenaceum* and *M. nivale* [116]. Similarly, mineral nitrogen applications increase the number of *F. culmorum* propagules [108]. However, in a two-year study, Teich (1989)[90] showed that nitrogen application limited the disease in the first year [99], although this decrease was not confirmed in the second year [100]. Fauzi and Paulitz (1994)[120] also demonstrated that disease levels were similar in the presence and absence of applications of 140 kg/ha ammonium nitrate.
There are two possible reasons for the uncertainties concerning the effect of nitrogen applications on the disease. Firstly, in certain cases, the applied nitrogen may be poorly distributed in the soil. *Fusarium* foot rot levels have been shown to decrease if ammoniacal nitrogen is applied with a syringe to the tilled layer but not if it is mechanically applied to the field [121]. Secondly, there may be an as yet unidentified link between attacks of *Fusarium* foot rot and head blight [6]. Thus, nitrogen-containing fertilisers may have a different effect on the *Fusarium* head blight and thus affect the production of head blight inoculum. This link may involve competition between the species of *Fusarium* more specific to foot rot (such as *F. solani*) and those more specific to head blight (such as the species of the *F. roseum* group). Indeed, whereas germination of the conidia of *F. solani* is decreased by nitrate application, spores of the species of the *F. roseum* group use all forms of nitrogen [122].

Different forms of nitrogen have different effects. Urea appears to reduce head blight levels more than ammonium [90,123]. Teich (1989)[90] suggested three hypotheses to account for this difference between urea and ammonium: i) the urea may decrease the size of the *Fusarium* population because nitrite blocks reproduction in these fungi and inhibits the formation of chlamydospores [124], ii) urea may prevent the maturation of *Fusarium* ascospores, as it does for those of *Venturia inequalis* [125], iii) urea may increase the number of actinomycetes in the soil [122,126], these microbes being antagonists of *Fusarium graminearum* [126] and *F. oxysporum* [122]. Furthermore, rotting of the base of the stem and of the roots is more severe if nitrogen is applied in an ammoniacal form than as nitrate [121,122,127]. This may also result from the higher rate of germination of *F. solani* (which may cause foot rot) in the presence of ammoniacal nitrogen than in the presence of nitrate [122].

In contrast to what was observed for bunt [103], the application of phosphorus has been shown to limit foot rot [128] and head blight [99]. However, the result obtained for head blight was not confirmed in the second year of the study [100].

Potassium application decreases transpiration by causing stomatal closure, thereby helping to decrease the humidity of the air. Thus, the application of potassium limits the germination of rust spores [103]. Furthermore, high levels of potassium favour the synthesis of high-molecular weight compounds (such as cellulose), decreasing the availability of nutrients to
the pathogens and increasing mechanical resistance to parasite penetration [103]. However, potassium applications do not appear to affect the incidence of head blight [100].

To understand the meaning of these contradictory results, we need to bear in mind that the effect of fertiliser applications depends on two factors: the age of the plant and the balance of mineral elements. Indeed, foot rot levels are decreased by early applications of nitrate but increased by late nitrate applications [128]. Conversely, foot rot levels are increased by early potassium applications and decreased by late potassium applications [128]. Moreover, the application of manures with a poor nutrient balance leads to more severe foot rot symptoms [14,64]. Antagonism between nitrogen and potassium is generally reported [14,103]: excess potassium and nitrogen deficiency render the plant more resistant whereas potassium deficiency and excess nitrogen render it more susceptible. This antagonism results from the fact that potassium increases nitrate absorption [122]. The nitrate absorbed activates nitrate reductase, leading to the accumulation of amino acids [127]. These amino acids are used in the synthesis of organic nitrogenous compounds [122] and of high-molecular weight compounds [103]. These compounds, like cellulose, contribute to the resistance of plants to diseases by increasing mechanical resistance to parasite penetration and by decreasing the quantities of soluble amino acids and low-molecular weight carbohydrates, which serve as nutrients for these pathogens [103]. Soluble amino acids and low-molecular weight carbohydrates are produced by increasing the activation of nitrate reductase in the presence of excess nitrate, by activating enzymes such as amylase, proteases and glucosidases, and by decreasing phosphorylation in conditions of potassium deficiency [103]. This increases the intracellular concentration of these compounds [103], favouring the development of pathogens.

Maize plants are most resistant to diseases when sugar concentrations are high [129]. Sugar levels are inversely proportional to nitrogen supply and directly proportional to potassium supply [14]. Conversely, if we consider three, rather than two, mineral elements, the mineral balance least favourable for the development of foot rot in wheat, both in seedlings and in adult plants, is high levels of nitrogen and potassium and low levels of phosphorus [128].

Several studies have demonstrated the existence of a relationship between the mineral and organic composition of the soil and the development of soil microorganisms, which may antagonise or stimulate the development of *Fusarium*. Thus, applications of organic fertiliser
rich in nitrogen, particularly the ammoniacal and nitrous acid forms, significantly reduce (by a factor of 1000) the size of the populations of many soil pathogens and markedly increase the size of populations of soil microorganisms [123]. Furthermore, a mixture of ammoniacal nitrogen and ammonium is recommended to increase the capacities of *Pseudomonas fluorescens* populations, thereby favouring the development of the plant (as this bacterium is a rhizobial bacterium) and inhibiting the development of *Fusarium* [130]. An application of glucose (simulating root exudates) to soils containing chlamydospores of *F. oxysporum* and *F. solani* favours the germination of these spores and the development of their germination tubes [131]. A similar result was also reported in a previous study: wheat root exudates increase the number and length of germination tubes arising from the spores of *F. roseum* and *M. nivale* [98]. The effect of glucose application is even greater if the soil is nutrient-poor, especially for slow-growing species such as those of the genus *Fusarium* [132].

The mineral nutrition of plants and the development of microorganisms in the soil also depend on soil pH. The conidia of *M. nivale* do not germinate in culture *in vitro* if the pH of the medium is lower than about 5 [11]. However, pH has been found not to affect the incidence of the disease [99,100].

2-5. **The effect of sowing date**

Sowing date is another element of crop management that has an indirect effect on the production of, and infection by spores, because it partly determines flowering date, together with the variety sown and climate. If the sowing date is such that flowering coincides with spore release, then more frequent and severe attacks are likely. This hypothesis is based on the fact that the anthesis is the stage in wheat at which the consequences of pathogen attack are thought to be the most severe [12,16,86,87].

The effect of sowing date should therefore be considered with respect to the level of attack as a function of the earliness of the variety. Early-maturing wheat cultivars tend to be more resistant to head blight than cultivars that mature later [133]. The duration of the growth period in wheat has a significant positive effect: the longer the cycle of the variety, the lower the degree of contamination observed [134]. As this effect is not observed in barley and oats, Couture [134] suggested that the threshold number of days exceeds the length of the cycle for wheat, but not for barley and oats. However, conflicting results were obtained in other studies: resistance to head blight has been shown to be independent of maturity factors [135], and late sowing has been shown to favour the development of head blight whereas early
sowing favoured the development of foot rot [16]. The results obtained almost certainly
depend on the climate of the region concerned and the variety grown. Flowering period and
the duration of the growth period for a given variety probably depend essentially on the year
and region concerned: the only valid way to determine the characteristics (period and duration
of flowering and growth duration) of varieties adapted to a given region is to carry out a
frequency analysis of the climate, which requires the monitoring of head blight attacks in a
given region over many years. Sowing on several dates (to extend the flowering period)
should make it possible to limit the risk of heavy contamination in all the fields. A frequency
analysis in a given region may also provide useful information for the optimal choice of
sowing date.

2-6. Effects of canopy density

If the inoculum is dispersed primarily by splashing, the density of the canopy is an important
factor as it may place obstacles in the way of this dispersal. Thus, a low canopy density can
favour spore dispersal (by creating fewer obstacles). In the case of high canopy density, due
to nitrogen fertilisation and/or higher sowing density and/or smaller spaces between rows of
wheat, two opposing effects may be observed. Firstly, the high density is likely to increase the
humidity of the canopy [116], favouring spore germination, as has been observed for rusts
[103]. Alternatively, the high density of the canopy may increase the number of obstacles,
limiting the vertical dispersal of spores towards the ear. However, for Fusarium infections,
more severe attacks of foot rot [16] and head blight [5] have been reported in cases of high
population density: more than 350 grains sown per m² [16].

Plant population density may also be increased by weed infestations. In the presence of a
large weed population, the number of species of Fusarium causing head blight in wheat
increases, as does the intensity of the disease [1]. This may be accounted for by the number of
common field weeds (both dicotyledonous and monocotyledonous weeds) on which
numerous species of Fusarium have been detected [62]. Table 2 lists the various weeds
known to serve as hosts for Fusarium, and the species of Fusarium found on them.

The "host-plant" nature of these weeds [14,48,62,98] suggests that they constitute a potential
source of inoculum. Since 1950, grass weeds have been suspected to act as a source of
inoculum whereas other weeds have been thought to favour infection in a different way, by modifying the microclimate, increasing heat and humidity [136].

Whatever the role played by the various weed families, the decrease in the incidence of the disease following the use of herbicides suggests that weeds do have an effect [99]. However, different results were obtained in the following year of the same experiment [100]. This difference probably results from interactions between cultural practices and weeds. For example, the withering of forage grasses (fescue, cocksfoot, ryegrass) due to *F. roseum* increases with the dose of nitrogen applied [14].

### 2-7. Effects of infestation of the canopy by bioaggressors others than fungi responsible for the head blight

The term "canopy" is often used to describe the green canopy, consisting of healthy plants. Diseases, particularly foliar diseases, decrease the area of the plants capturing light, thereby limiting the development of the crop. Very few studies have focused on the competition between pathogens or between diseases. Only two authors have analysed the effect of a powdery mildew attack on head blight, and the results of these two studies conflicted. In the first study, powdery mildew attacks were found to have a positive effect on head blight attacks [30] whereas in the second, no effect was observed [99]. These conflicting results may be accounted for by several factors, such as climatic conditions, the varieties cultivated and the intensity of the diseases in these studies.

In addition to soil microorganisms (discussed in section 2-2), the ear microorganisms play a role, notably in biological control [137]. Indeed, the plant may be protected against pathogen attacks by prior inoculation with pathogen isolates of various degrees of virulence, which is known to induce resistance in cases in which protection requires activation of the host's defence responses [138]. Thus, inoculation with *Microsphaeropsis* spp. significantly decreases the production of *F. graminearum* ascospores on wheat and maize ears and, to a lesser extent, on wheat and maize residues [137,139]. This decrease is particularly large if the residues are inoculated early with *Microsphaeropsis* (test with inoculation two weeks before the *F. graminearum* attack and 4 to 6 weeks after it) [137]. Prior inoculation with *Phoma betae* or *Pythium ultimum* decreases the severity of head blight on wheat and extends the incubation and latent periods of the disease due to *F. culmorum, F. avenaceum, F. poae* and *M. nivale* [138]. Such prior inoculation also significantly increases the number of grains per
ear and decreases the number of *Fusarium* isolates obtained from the harvested grains [138]. A previous study [140] showed that contact between *Pythium oligandrum* and *F. culmorum* macroconidia led to coagulation and a loss of cytoplasm, followed by complete degradation of the walls of the macroconidia and the production of *Pythium oligandrum* oogonia in abundance on the parasitised macroconidia. Inoculation of wheat ears with *Alternaria alternata, Botrytis cinerea* or *Cladosporium herbarum* at GS69 (Zadoks growth scale, Zadoks et al., 1974[141]) favours the infection of ears inoculated with *F. culmorum* at GS65, whereas prior inoculation, at GS59, decreases infection of the ears by *F. culmorum* [142]. Following the demonstration of antagonism between these saprophytes, Liggitt et al. [142] suggested that *Alternaria alternata, Botrytis cinerea* and *Cladosporium herbarum* produce volatile antibiotic substances that limit the growth of *F. culmorum*, consistent with the decrease in size of *F. culmorum* colonies observed following prior inoculation with other saprophytes.

Such antagonism is also observed between *Fusarium* species [22,27] and between species of *Fusarium* and *Microdochium* [22]. This is the case, in particular, for fungicide treatments acting against only one of these genera, resulting in the development of the unaffected genus to a greater extent than in the absence of treatment, when the two genera compete [21,22]. Similarly, prior inoculation with germination fluid from *F. avenaceum* predisposes wheat ears to infection by conidia of *F. avenaceum* or *M. nivale*, but not by conidia of *F. culmorum*, or *F. poae*. Other germination fluids were found to have no such effect [138]. This suggests that diffuse substances present in the germination fluid of *F. avenaceum* promote the production in competitive host tissues of resistance factors specific for *F. avenaceum* and *M. nivale* [138].

2-8. Effects of chemical treatments

The efficacy of fungicide treatments for head blight is variable, and difficult to predict. It depends not only on the active ingredient but also on the method and date of application of the fungicide [6,27]. If a fungicide treatment is to be effective, it must be applied several days before the attack and entirely cover all the ears [142]. Efficacy also depends on the interaction between fungal development and climate [143], the virulence of the pathogenic strains [143] and above all, the parasite complex present on the plot of land concerned [6]. Each of the fungal species implicated in head blight has a specific fungicide susceptibility profile: *F. graminearum* is particularly susceptible to triazoles whereas *F. avenaceum* is more susceptible to strobilurins. *F. roseum*, a complex consisting of *F. graminearum, F. culmorum* and *F. avenaceum*, is much more susceptible to triazoles than is *M. nivale*, which is
essentially susceptible to strobilurins [22]. Triazole fungicides containing tebuconazole are currently the most effective [21,27,143].

Chemical treatments may have an effect by modifying the height of the canopy. For a single variety grown in a single year, in a single region, the use of plant growth regulators (primarily gibberellin inhibitors) results in more severe head blight attacks [14,64,118,120]. This is presumably because plant growth regulator treatment results in the ears being closer to the soil, and therefore to the crop residues (source of inoculum), facilitating the dispersal of the spores to the ear (by splashing in particular). Indeed, the distance of the ear from the soil is a factor for resistance by avoidance in cultivars. Plant growth regulators also affect the microclimate of the crop residues on the soil [120], possibly increasing the production of perithecia and ascospores.

3- Effects of various cultural practices on the production of mycotoxins by *Fusarium*

The effects of cultural practices on the intensity and severity of head blight symptoms are well documented, at least for certain practices, but far fewer studies have considered the effects of these practices on mycotoxin production.

3-1. Mycotoxins produced by *Fusarium*

Fungi of the genus *Microdochium* are currently considered not to produce toxins [8] and will therefore not be dealt with in this section. It should also be noted that studies on the mycotoxins produced by *Fusarium* have generally focused on deoxynivalenol (DON), even though *Fusarium* also produces other types of mycotoxin. Figure 2 presents the chemical structures of the main mycotoxins produced by *Fusarium*.

Deoxynivalenol belongs to the trichothecene family, which contains two types of toxin: type A and type B. Type A toxins include toxins T2, HT2, diacetoxy-scirpenol (DAS) and scirpenol [39], which act on the initiation of protein elongation [144]. Type B corresponds to deoxynivalenol (DON or vomitoxin) and its acetyl derivatives — 3-acetyldeoxynivalenol (3acDON) and 14-O-acetylDON-4 — nivalenol (NIV) and verrucarol [39]. The toxins of this group act on the elongation and termination steps of protein synthesis [144]. Trichothecenes are toxic to all organisms. In mammals, the absorption of trichothecenes leads to blood and
digestive disorders. In humans [145], burning sensations in the mouth and stomach, headaches, a decrease in red blood cell count, bleeding, necrosis of the throat and stomach and, in some cases, death, were observed in a humanitarian catastrophe in the USSR between 1942 and 1947 caused by the ingestion of wheat that had suffered foot rot all winter [38,39]. The production of trichothecenes is favoured by cold and humidity [146] and studies on various organisms have demonstrated the toxicity of these substances. In animals, vomiting is observed after the ingestion and absorption of at least 10 mg deoxynivalenol per kg, together with irritation of the mucous membrane lining the mouth and the oesophagus, with animals refusing to eat after ingesting 2 mg deoxynivalenol per kg.

In plants, deoxynivalenol retards the germination and growth of wheat, and inhibits the growth of the grain and the coleoptile tissues [147]. The phytotoxicity of mycotoxins is generally estimated by means of coleoptile elongation tests [43,145]. In adult plants, deoxynivalenol seems to circulate in the phloem, with the concentration of this molecule in the plant following a descending gradient from the rachis, through lemmas and grains to the peduncle [148]. In addition, from the fourth day after inoculation, the flower parts, rachis and peduncle contain larger amounts of deoxynivalenol below the point of infection than above it [149].

Fusarium can also produce mycotoxins of two other families: fumonisins, which are recognised carcinogens [13,150] and zearalenone. Zearalenone belongs to the oestrogen family. It perturbs reproductive hormone equilibrium and alters secondary sexual characters [145]. Pigs are particularly sensitive to this toxin [145]. In the laboratory, it has been shown that this compound also controls reproduction in the fungus that produces it, by regulating peritheciun production [151-153]. No phytotoxic effect of zearalenone has been observed in coleoptile elongation tests [145].

To account for the small number of studies dealing with the mycotoxins produced by Fusarium on wheat, it should be borne in mind that most of these studies have been carried out in Europe and North America, where type A trichothecenes are rarely detected [154]. Furthermore, fumonisins and zearalenone are found essentially on maize [8,155]. In addition, zearalenone is considered to be only mildly toxic to humans [155].
3-2. Effect of soil tillage

Soil tillage seems to have similar effects on the production of mycotoxins and on the disease itself (see part 2-1). Unlike chisel ploughing and direct drilling systems, mouldboard ploughing decreases the concentration of deoxynivalenol in the grains [19,101,156]. In one study, a significant positive correlation between deoxynivalenol concentration and the intensity of soil tillage (ploughing or direct drilling) after a maize crop was observed [100]. Similarly, another study comparing three cropping systems involving ploughing (productive, integrated and organic) with a direct drilling system over two years reported that levels of deoxynivalenol, zearalenone and nivalenol contamination were highest for the direct drilling system [55].

3-3. Effect of the preceding crop

The effect of the preceding crop is also similar for disease symptoms and deoxynivalenol concentration. Deoxynivalenol concentration in wheat is 25% lower after a soybean crop than after a wheat crop and 49% lower after soybean than after maize [19]. Wheat following maize in the rotation has been found to have deoxynivalenol concentrations six times higher than those in wheat following another cereal (wheat, barley) or soybean [100]. In direct drilling systems, wheat crops following an oilseed rape crop contain 90% less deoxynivalenol than wheat crops following a maize crop [101]. As for soil tillage, only few studies have produced useful results.

3-4. Effect of mineral nutrition

We saw in section 2-3 that the effects of mineral nutrition on the disease are unclear. In contrast, there are fewer strong contradictions concerning the effect of mineral nutrition on mycotoxin production. This is probably because very few studies have focused on this question. Increases in nitrogen input from 0 to 80 kg/ha, result in increases in grain deoxynivalenol content; at levels above 80 kg/ha, a small but significant decrease was observed [119]. Urea-based fertilisation resulted in lower levels of deoxynivalenol in wheat grains than did ammonium nitrate in 1997, but this result was not observed the preceding year, in which no effect had been demonstrated [156]. Applications of ammonium-nitrate-urea solution and nitramoncal did not give significantly different results [119]. Potassium, phosphorus and pH did not appear to affect deoxynivalenol concentration significantly [100]. A laboratory study showed that Mg\(^{2+}\) ions inhibit the elements responsible for inducing trichothecene synthesis [38].
These studies were empirical, and were not based on any particular assumption concerning the mechanisms involved. It therefore remains difficult to extrapolate the results obtained.

3-5. Effect of variety earliness

Very few studies have focused on the effect of wheat variety earliness on the production of mycotoxins. The earliest varieties seem to accumulate more deoxynivalenol than do late-flowering varieties [157], possibly due to varietal differences but more probably due to greater coincidence of the phase of maximum susceptibility in plants with the period most favourable for spore dispersion. This trend appears to be based on the relationship between the severity of the attack and the intensity of toxin contamination. The choice of variety is therefore of key importance for combating the accumulation of toxins in the grains [158].

3-6. Effect of fungicide treatments

The application of a triazole fungicide, such as tebuconazole, decreases levels of deoxynivalenol contamination in artificially inoculated crops with respect to those observed in untreated controls [21,27,28,143]. However, conflicting results have been obtained [22]. More generally, in conditions of natural contamination, the application of fungicides does not significantly decrease the concentrations of deoxynivalenol and nivalenol and there is not necessarily a correlation between the application of fungicides and the quantity of deoxynivalenol and/or nivalenol found in the grains [6,55]. This may be accounted for by the multitude of species present in fields and the relative specificity of fungicide treatments for only one or a few species. Another hypothesis has also been put forward: too low a dose of fungicide may stimulate the production of deoxynivalenol by *Fusarium* in wheat grains [53].

This analysis shows not only that few studies have investigated the effects of cropping systems, but also that the characterisation of the disease in these studies is often severely flawed. This makes it difficult to extrapolate and to interpret the results. To understand the relationship between cropping systems and mycotoxin production, we therefore need to break down this relationship into two parts: the effects of cropping systems on the disease (symptoms, nature of the pathogenic agent) (section 2) and the relationship between the disease and the production of mycotoxins, which may itself vary as a function of cropping system.
4- Relationship between the disease and toxin production

4-1. Toxin production by the fungi

Toxin production in fungi is controlled by many factors. One of the most important factors intrinsic to the fungus is the genetic capacity of the pathogenic strain to produce toxins, in terms of the quantity and type of toxins produced [34,159,160,161]. Thus, strains of *F. graminearum* may, like *F. culmorum*, produce isomers of monoacetyldeoxynivalenol or alternatively, like *F. crookwellense*, may produce acetyl derivatives of nivalenol [4]. Strains of *F. sambucinum* may produce the T2 toxin whereas those of *F. sporotrichioides* produce diacetoxyscirpenol (DAS) [4]. It should be noted that strains of *F. graminearum* [60,159,162] and of *F. culmorum* [163,164] tend to produce either deoxynivalenol or nivalenol, but not both. Thus, the two toxins are generally not produced by the same strains, although the two types of strain often coexist in the same field [165]. Moreover, regionalisation of the types of strain at world level is observed for *F. graminearum* strains producing 3-acetyldeoxynivalenol and those producing 15-acetyldeoxynivalenol, with strains producing 3-acetyldeoxynivalenol predominating in Europe, China, Australia and New Zealand whereas those producing 15-acetyldeoxynivalenol predominate in the United States [159]. Data for a collection of 188 strains of *F. graminearum* indicated that European and American strains produce essentially deoxynivalenol and are more aggressive than Nepalese strains, which produce either deoxynivalenol or nivalenol [166]. A negative correlation has been found between the production of zearalenone and that of trichothecenes (deoxynivalenol or nivalenol) by the spores of *F. graminearum* [52]. This appears to indicate that the production of these toxins shares a common control process, which may play an important role in the ecology of *F. graminearum* and *F. pseudograminearum* [52].

Other factors may also cause variations in toxin formation. These factors include the substrate, the period of colonisation of the substrate by the strain, competition between microorganisms [2,34,52,53], interaction between the pathogenicity of the strains and the wheat variety [167,168] overall climatic conditions [51] and, more particularly, temperature [2,52-54], humidity [2,52,53] and rainfall [54].
Other poorly understood factors extrinsic to the fungus also operate. For example, the protein to carbohydrate ratio, which is important in mycotoxin synthesis [169], may play a role. However, Bakan (1998) [38] showed that toxin formation is not necessarily controlled by food stresses and is not necessarily linked to the growth of the fungus.

This last point is particularly important because many authors have suggested that the trichothecenes (deoxynivalenol in particular) produced by *Fusarium* species play a role in the aggressiveness or virulence of certain isolates [164,170-174]. Indeed, certain mutants deficient in deoxynivalenol are less aggressive on wheat than the original strain [175]. Similarly, the least virulent mutants of *F. graminearum* in terms of plant colonisation are those unable to produce deoxynivalenol or nivalenol (mutations in the Tri5 gene, involved in trichothecene production) [171]. Thus, trichothecenes are considered to be virulence factors in *F. graminearum*, during the infection of wheat [145,175], and in *F. culmorum*, during the infection of barley [164].

In conditions of artificial contamination with a given pathogenic strain or in natural contamination conditions, several authors [19,27,90,100,147,164,176,177] have observed a significant, positive correlation between the incidence and/or severity of the disease and deoxynivalenol concentration (also reported by Bai *et al.*, 2001 [49] for 116 wheat lines). Others [26,49,178,179] have demonstrated a significant correlation between deoxynivalenol concentration and the fungal biomass of the grains (measured by assessing the amount of ergosterol). These results suggest that new cultivars could be selected on the basis of symptoms to ensure low levels of deoxynivalenol. However, exceptions were found among these lines: certain cultivars present severe symptoms with low deoxynivalenol levels, particularly those with moderate resistance to propagation of the pathogenic agent [49]. Thus, the production of deoxynivalenol is not essential for the infection of grains [164]. Furthermore, after the inoculation of five wheat varieties with a strain of *F. graminearum*, no correlation was observed between deoxynivalenol concentration and the severity of infection [172]. Similarly, after inoculation with a complex of species, no correlation was observed between deoxynivalenol concentration and infection on wheat, barley or oats [178]. In conditions of natural contamination, no correlation has been found between the intensity of the disease on wheat and the concentration of deoxynivalenol [53,55,78], or between the presence of *Fusarium* and the concentration of deoxynivalenol [154]. This lack of relationship is also observed with other toxins, such as nivalenol [55]. A similar result was obtained with
maize: after inoculation of the ear with *F. poae*, no relationship was found between the severity of the disease and toxin (nivalenol and fusarenon X) production [180]. These differences in results may be due to the ranges of variation tested and the comparisons made not being the same. Some comparisons were made in the field and others in the laboratory; some were made after artificial inoculation with one or several species and others were made in natural contamination conditions.

The date of the attack and the species involved also affects the relationship between symptoms and toxin levels. Indeed, it is only to be expected that as the number of affected grains increases, so does the risk of major contamination by mycotoxins. However, a late attack may prevent the disease from reaching a high level of severity, even if very high levels of mycotoxins are produced. Thus, fungi and mycotoxins are detected before any visual symptoms of disease [148,181]. In addition, not all the species of the parasite complex responsible for head blight produce mycotoxins: *Microdochium*, for example, does not produce mycotoxins. Similarly, even among *Fusarium* species potentially capable of producing toxins, strains unable to produce toxins have been identified [2,38,162,182]. Other strains have been identified that produce far more toxin than the mean amount for all strains [183]. In addition to climate (particularly rainfall levels and temperature before and at flowering), the species profile of the fungi responsible for head blight also depends on agronomic factors such as soil tillage, nitrogen fertilisation, fungicide use, crop rotation and host genotypes [10]. Furthermore, the species of *Fusarium* potentially able to produce toxins cannot necessarily produce trichothecenes (the most studied mycotoxins) [183]. These observations indicate that it is possible for the plant to present symptoms in the absence of toxin production. However, the conditions of toxin productions in the field remain mostly unknown.

The relationship between the intensity or severity of symptoms and toxin production also depends heavily on the type of resistance carried by the plant. The severity of symptoms is known to depend heavily on plant resistance: certain genotypes limit the development of the mycelium in the grain, protecting the grain against degradation and limiting the visual signs of attack, but are not very tolerant of mycotoxins, with very high mycotoxin levels being recorded [49]. Conversely, other types of cultivar may present severe symptoms with only low mycotoxin levels [49].
4-2. Different mechanisms of genetic resistance to *Fusarium* head blight in plants

**Non-specific resistance**

It is not easy to identify cultivar resistances because the species of *Fusarium* responsible for head blight are saprophytes with a broad host-range and low levels of intraspecific variation, which is not the case for pathogens in general [6,18]. It should also be noted that all cultivars are susceptible to some extent, with even the least susceptible cultivars displaying only moderate resistance [16,49,50]. Resistance to the development of the fungus and to the accumulation of trichothecenes is probably controlled by different genes [43].

**Heritability of the resistance**

The low specificity of *Fusarium* and *Microdochium* generates polygenic resistance in the plant, involving genes with various degrees of dominance [43,184]. It is the combination of these genes, controlled by the environment [176] that results in genetic resistance in the plant [13].

The inheritance of resistance can be described by a dominance-additive effect model, in which the additive effect is the essential factor (results based on diallel analyses) [9,25,185]. Several studies [18,25,186-188] have demonstrated the existence of resistance genes on 18 of the 21 chromosomes (1B, 2A, 3, 3B, 4B, 5A, 6B, 6D, 7A, 7B etc.). Three pools of genes are responsible for three different genetic constitutions conferring different types of resistance to head blight in wheat: in Eastern European winter wheats, in Japanese and Chinese spring wheats and in Brazilian and Italian spring wheats [189]. The most commonly used source of resistance is undoubtedly Sumai 3 (in China), in which stable resistance is combined with valuable agronomic characteristics (yield potential and resistance to rusts and powdery mildew) [18,43,190]. The resistance of this genotype, like that of Frontana (a Brazilian variety) and Nobeokabouzu-komugi (a Japanese variety) — the two other major sources of resistance — is based on two or three genes with additive effects, the sensitivity of which varies with a multitude of minor genes [9,18,191]. The crossing of Frontana and Sumai 3 resulted in the detection of two major quantitative trait loci (QTLs) [13]. These sources of resistance are of great potential value for future crosses and the selection of new cultivars because they are stable.

Resistance to head blight in wheat is horizontal (non-specific) and thus protects against all species of *Fusarium* and *Microdochium* [6,18,43,192].
No correlation has been found between seedling resistance and resistance in the ear. Thus, a given cultivar may be susceptible at the seedling stage and resistant at the adult stage or vice versa. Alternatively, it may be susceptible throughout the crop cycle [5]. The activation of resistance genes during an attack depends on the developmental stage of the wheat, varying throughout the crop cycle [5,47,49,50,90]. Peak toxin accumulation also depends on the cultivar grown [193]. Finally, it should be noted that the duration of the induction period and the intensity of the response to the defence mechanisms induced also determine the defence of the plant [43]. Tetraploid wheats are more resistance to head blight than are diploid wheats [194].

The defence or resistance mechanisms of the plant depend not only on growth stage, but also on the humidity and temperature of the air [47] and on the genetic capacity of the wheat cultivar [5,18,49,50,90]. However, no plant host defence reaction has ever been detected macroscopically or histologically on the outside of the glumes [12]. In addition, no cellular or anatomical characteristics [13,50] or histological features [50] have been found to be associated with resistance or susceptibility. However, the techniques currently used, and ultrastructural studies in particular, have shown that susceptible and resistant wheat varieties react differently to infection and to the propagation of pathogens [94].

This implies the existence of multiple mechanisms of defence or resistance in plants, both active (including physiological processes) and passive (including morphological characteristics such as avoidance), and/or tolerance [5]. Five types of cultivar resistance are currently known and described.

*Types of resistance*

**Type I: Resistance to initial infection** [50]. Many cultivars may be resistant to initial infection. According to Nakagawa (1955) [195], this type of resistance is controlled by dominant genes at three loci subject to epistasis. This type of resistance may be passive or active.

*Passive mechanisms* involve morphological characteristics facilitating avoidance, making it possible for the plant to decrease the severity of the disease. The receptiveness of cultivars to the disease is lower in cultivars with awns [5,14,134], regardless of varietal differences in
resistance [134]. Similar results have been obtained with wheat, barley [134] and oats [45,134]. Similarly, the height of the ear and its angle with respect to the stem are negatively correlated with the severity of head blight and the accumulation of deoxynivalenol [191]. The receptiveness of cultivars to *Fusarium* and/or *Microdochium* is also lower for ears with a large peduncle — at least 15 cm between the flag leaf and the ear [5] — and/or without growth arrest, reducing the area in contact with the conidia and the duration of grain humidity [5].

A cultivar well adapted to its environment is also a criterion for resistance to head blight [18,43]. Indeed, a canopy with too many ears [5,16] or a flowering season that lasts too long [2,90] is likely to result in slightly higher susceptibility to the disease. The resistance of a variety may be evaluated as a function of the duration of time for which the flowers remain open and of the percentage of flowers presenting autogamous flowering (unopened anthers) and allogamous flowering (opened anthers) [50]. Studies have shown either that resistance to head blight may be maximised by the use of early varieties [133] or that earliness has no importance [135]. However, the importance of this factor seems to depend on the date of the attack and, thus, on the year and region studied. Similarly, only one study has investigated the effect of flowering date on mycotoxin contamination: the time of infection by *F. culmorum* that produces the greatest amount of deoxynivalenol contamination is restricted to a short period during anthesis [87]. However, like disease, flowering period and variety growth duration probably depend essentially on the year and region concerned: the only valid way to determine the characteristics (period and duration of flowering and growth duration) of varieties adapted to a given region is to carry out frequency analysis of the climate, which requires the monitoring of mycotoxin contamination in a given region over many years.

It is widely accepted that morphological characteristics are less important than the possible physiological resistance of cultivars [9]. This physiological resistance includes all the other possible mechanisms. *Active mechanisms* include defence reactions concerning the physiological qualities of the cell that limit colonisation of the plant by the fungus, such as the activation of enzymes degrading the fungal cell wall. Such a mechanism has been proposed for pathogenesis-related proteins (PR proteins) and hordeithionins [196]. Mechanisms of this type may account for the induction of several defence genes 48 hours after inoculation, including genes encoding chitinases, glucanases, peroxidases and thaumatin-like proteins (TLP) [197,198].
The endo- and exochitinases in plant cell walls help to improve plant resistance by hydrolysing chitin, a protective polysaccharide present in the cell walls of the pathogens [18,199]. The various types of chitinase known do not have the same effects [200]: the class I chitinase-a of rice strongly inhibits re-extension of the hyphae and is present in mature cells in particular. It collects at the tip of the hyphae, on the lateral walls and in the septa and has a very high affinity for fungal cell walls. This enzyme releases a large amount of reducing sugar from the fungal cell walls. It also ligates the lateral walls and the septa, is part of the walls of mature cells and degrades mature chitin fibres. In contrast, the class II chitinase-c of rice is found essentially at the tip of the hyphae, which it ligates before degrading young chitin fibres. Thus, chitinase-a is more effective than chitinase-c at inhibiting the growth of the fungus [200].

The antimicrobial activity of thionins is attributed to their capacity to create pores in the membranes of fungi by means of interactions between the phospholipids and phosphoinositides of the membrane or by decreasing the activity of enzymes by reducing disulphide bonds [199].

Thaumatin-like proteins may have two modes of action: they may be specifically produced in response to the presence of pathogenic fungi in a certain number of plants, including wheat [201] or they may disturb the signal transduction cascade in the cell, increasing tolerance to trichothecenes and favouring the development of fungi in the plant [198]. A trypsin-like protease produced by *F. culmorum* may be one of the key enzymes in the colonisation of plants by the fungus [202]. Indeed, the presence of this enzyme in barley has been shown to be correlated with the degradation of specific buffer proteins in infected grains [203]. The alkaline protease produced by *F. culmorum* is also involved in the colonisation of barley and wheat grains [204]. Peroxidases (POX) are oxidoreductive enzymes involved in the construction of the cell walls of plant hosts (phenol oxidation, suberisation, lignification) during defence reactions against pathogenic agents [205]. The concentration of POX increases significantly during the milk stage in wheat anthers inoculated with *F. graminearum*, whereas this is not the case in healthy wheat [205].

Polyphenol oxidases (PPO) are involved in the oxidation of quinone polyphenols (a microbial component) and in the lignification of plant cells during microbial invasion [205].
specific activity of PPO is maximal during the milk stage in wheat and declines rapidly thereafter [205]. Levels of PPO activity are higher in resistant than in susceptible varieties and increase following inoculation with *F. graminearum* [205].

One possible strategies for reducing the risk of mycotoxin contaminations is the development of more resistant transgenic crops. Three approaches are possible [206]: (i) transgene-mediated control of the ability of *Fusarium* to infect and colonise the ear, through the overproduction of specific antifungal proteins and metabolites, or by increasing the plant’s own defense systems in kernel tissues; (ii) the prevention of mycotoxin biosynthesis, or the detoxification of mycotoxins in plants and (iii) the development of more resistant plants based on transgenic crops engineered to produce a *Bacillus thuringiensis* (Bt) toxin. Some Bt maize hybrids have the potential to reduce the level of fumonisin B produced by *F. verticillioides* [206]. This effect probably results from a decrease in sensitivity to one of the possible means of ear tissue contamination: attack by insects [206,207]. Indeed, *Bacillus thuringiensis* (BT) is known to produce proteins toxic to insects [208]. This strategy has not yet been applied to wheat.

Type I resistance is generally estimated by determining the percentage of spikelets infected seven to 21 days after inoculation or at maturity [9]. However, the number of infected spikelets does not necessarily reflect the total damage caused by the pathogen. The symptoms also depend on the severity and rapidity of rachis invasion by the pathogen.

**Type II: resistance to (kinetic) propagation of the pathogenic agent** in the tissues [50].

The mechanisms involved in this type of resistance are purely active, such as inhibition of the translocation of deoxynivalenol [18] by an ABC (ATP-binding cassette) transporter protein, the Pdr5p (pleiotropic drug resistance) [209] or increasing the stability of cell membranes [18]. This process limits the propagation of *Fusarium culmorum* and *F. graminearum* [95] from the glumes [18,26] and/or ovaries and glumes to the rachis and the pedicel [94-96]. The endo- or exocellular migration [94-96] of the fungus leads to a series of changes in host cells, including degeneration of the cytoplasm, organelles (e.g. chloroplasts) and cell wall [95,96] by enzymes such as cellulases, xylanases and pectinases [95,96] produced during the penetration and colonisation of the tissues of the wheat ear [95,96]. Once attacked, susceptible varieties respond to infection only by forming a very thin wall in the periplasmic space of the infected tissues [94]. In contrast, resistant varieties react strongly: extensive formation of a
very thick wall located close to the infected cells and of a large papilla formed by the rapid
and intense deposition of callose (β-1,3-glucan), lignin and other compounds [94]. Three days
after inoculation, defence structures are more solid, with denser cell walls, particularly in cells
adjoining contaminated cells, in the envelopes of cultivar Frontana (resistant) than in those of
cultivar Agent (susceptible) [210]. The process of lignification may be involved in plant
defences, establishing mechanical barriers to pathogen invasion [94,210], modifying the
structure of cell walls and making them more resistant to the degradative enzymes produced
by the pathogen [210]. Furthermore, lignification may inhibit or reduce the diffusion of small
molecules or ions, including the mycotoxins secreted by the fungal hyphae in the host cells
[94,210], and reduce the movement of nutrients from the host cell to the pathogen [210]. In
addition to the lower permeability of the cell membranes in resistant genotypes, the basis of
the inhibition of deoxynivalenol translocation may also involve a lower affinity of the
membrane for deoxynivalenol [26]. This hypothesis is based largely on observations in two
types of mutant yeast [144]. One of these types of mutant yeast presents a low affinity and
low cell membrane permeability for trichothecenes, and in the other, the 60S ribosomal
subunit has a lower than normal affinity for these toxins. The target of deoxynivalenol is a
cytoplasmic peptidyl transferase [211]. Mammalian and fungal cells have a modified peptidyl
transferase that is tolerant to trichothecenes [211]. In cases of type II resistance, the
trichothecenes seem to bind to the ribosomal 60S subunit, by methylation, thereby blocking
the translation of the RNA and inhibiting protein synthesis [38,43,145]. This inhibition leads
to the inhibition of peptidyl transferase activity, opposing resistance to trichothecenes. In
cases in which the membrane is not permeable to deoxynivalenol, a virulence factor
promoting fungal growth [26] the propagation of the pathogen agent is restricted. Thus, three
to five days after inoculation, susceptible varieties display more extensive propagation of the
fungus, higher levels of deoxynivalenol accumulation and fewer pathological changes in
infected tissues [94]. A hypersensitive reaction is also possible [18].

This type of resistance can be identified visually by observing the symptoms, which reflect
the propagation of the disease from the point of infection [9]. It is also characterised by low
levels of ergosterol (an indicator of fungal biomass) [3]. The most well known variety
displaying this type of resistance is Sumai 3 [189].

Type I and II resistances may be combined in a given genotype [18,50]. This results in a
variety tolerating the invasion of grains that displays no symptoms on the grains [50].
Similarly, the number of infected spikelets does not necessarily reflect the total amount of damage done by the pathogen. The amount of damage also depends on the severity and rapidity of the invasion of the rachis by the fungus. Thus, the three principal sources of variation in the expression of symptoms are the environment [57], the species and/or strain of *Fusarium* [57,212] and factors involved in the maturation of cereals that also play an important role in the epidemiological profile of the disease [212]. Wheat contamination is favoured by high levels of precipitation in the ten days preceding grain maturity, but only if the minimum temperature in the ten days following ear emergence is sufficiently high [212]. It should be noted that these factors are varietal and also depend on the fungal species concerned [212].

It seems likely that, in certain cases, the production of trichothecenes (deoxynivalenol in particular) by fungi activates their development (hyphal growth: type II resistance) and colonisation (type I resistance), by interfering with the defence response of the plants [18,26,175,213]. The quantity of trichothecenes in the grains is controlled by the degradation of deoxynivalenol and by the tolerance of the plants to this toxin [18].

Two other types of resistance have recently been discovered. Both involve defence reactions directed against trichothecenes, in which the synthesis of this toxin is blocked or its degradation enhanced [147,214].

**Type III: resistance involving the capacity to degrade** deoxynivalenol [214-216].

In some varieties (e.g. Fredrick), enzymatic systems for the detoxification (degradation) of trichothecenes, involving the acetyltransferase Ayt1p [209] for example, have been identified based on a decrease in the amount of deoxynivalenol six weeks after inoculation [215]. Deoxynivalenol begins to accumulate three days after inoculation (four days according to Savard, 2000[149]) of an ear. The concentration of this toxin increases, peaking after six weeks, and then decreases naturally to a concentration that remains constant thereafter at maturity, harvest and during grain storage [90].

This resistance was identified by inoculating grains of wheat, rice and triticale with a single strain of *F. graminearum* [214]. This study showed that the varieties most resistant to pathogen development (low *F. graminearum* biomass, as estimated from ergosterol content) have a higher ergosterol/deoxynivalenol ratio than do varieties more susceptible to the
development of this pathogen. This implies that the susceptible varieties are able to degrade the deoxynivalenol produced by the large number of fungi present.

This type III resistance, characterised by a high ergosterol/deoxynivalenol ratio [213], protects against symptoms due to the propagation of the fungus. However, there may nonetheless be a decrease in the number or weight of grains and their size [9]. This resistance is estimated visually by measuring yield and by comparing the values obtained for ears with and without symptoms [9] but this estimation also depends on resistance to grain infection.

**Type IV: resistance involving tolerance to high concentrations** of deoxynivalenol (apparent insensitivity to trichothecenes) [3].

Like cultivars with type II resistance linked to the modification of peptidyl transferase, the target of deoxynivalenol, cultivars with type IV resistance also have a modified peptidyl transferase [3]. This enzyme stabilises the membrane, conferring greater tolerance to trichothecenes [43]. It has also been suggested that changes in the permeability of the cell wall or in the signal transduction cascade in the cell inducing greater tolerance to trichothecenes by thaumatin-like proteins can account for this type of resistance [198].

For a given severity of disease, this resistance is generally estimated by a higher level of deoxynivalenol compared to other genotypes [9].

**Type V: resistance to grain infection** [5].

This type of resistance results in differences in yield despite similar levels of attack or, conversely, little effect on yield despite the evident presence of the disease after artificial inoculation [5,9,217]. In contrast to the other types of resistance, this type of resistance may not correspond to a physiological process, but rather to the “morphological” expression of an interaction between various complex processes: resistance of types II, III and IV.

4-3. **Resistance to head blight according to the composition of plant tissues**

In addition to genetic resistance, the composition of plant tissues is also a criterion of resistance to the development and/or propagation of the fungus and its toxins. The susceptibility of wheat cultivars is linked to the concentration of choline in the ear at anthesis [218], superoxide dismutase activity in the ears in cases of contamination by deoxynivalenol or *F. graminearum* [219] and the concentration of p-coumaric acid, one of the phenolic acid
precursors of lignin [210]. The quantity of this compound in the ear is much higher after inoculation of the glumes, envelopes and rachis with *F. culmorum* than in uninoculated tissues [210]: this is thus a very important factor determining the susceptibility of crops. The accumulation of an acetyltransferase encoded by *FsTri101* in the endosperm and glumes of wheat confers partial protection against *F. graminearum* [220]. Similarly, feluric acid in the ears seems to be involved in resistance to ear blight in maize [221], although healthy and inoculated ears contain similar amounts of this compound [210].

The composition of wheat and barley grains may also affect contamination levels [134]. Couture [134] suggested that a high gluten content in hard wheats and a high starch/protein ratio in malting barley could protect the crop against head blight.

The relative sugar content of leaves or stems has been proposed as an indicator of susceptibility to diseases [222]. Unlike powdery mildew, head blight is favoured by a low sugar content of the tissues [222]. Similar results were obtained in studies of stem blight in maize, involving a parasitic complex consisting of 70% *Fusarium graminearum* [129]. The refractive index of the sap of a section of pressed stem is used to assess the sugar content of stem bases (Bertrand’s method) and it has been shown that the higher the sugar content of maize stems, the more resistant to stem blight the maize is likely to be [129]. Messiaen [129] also indicated that *F. graminearum* is equally able to use sugars, starch and cellulose as sources of carbon, and proteins, ammonium salts or nitrates as sources of nitrogen. Thus, *Fusarium* makes use of storage proteins, and degrades the cell walls and starch grains of infected grains [41,42].

**Conclusion**

Head blight is a widespread disease with major consequences for health. However, it has been little studied in the open field. Indeed, field studies of the epidemiological cycle of the disease are hindered by a major problem: identification of the species and strains making up the parasitic complex of the inoculum. A mean of five to seven species, and up to nine in total [8], may be present in the same field in the same year. These species can only be identified after culture in Petri dishes and the observation of a collection of morphological criteria (colour, macroconidia, microconidia, ascospores, chlamydospores of a strain), which requires
the conservation of living spores, or molecular genotyping (which remains difficult for these species).

In the laboratory, the various steps in the plant infection process and in mycotoxin production are essentially studied from an environmental viewpoint. Uncertainties remain concerning, in particular, the necessity of inducing the development of perithecia by exposure to ultraviolet light, the natural climatic conditions in which spores are released (macroconidia and ascospores) and the sensitivity of these mechanisms to environmental factors.

In the field, spore dispersal depends on climatic factors in two ways: climatic factors affect spore production (see section 1-2) and strongly regulate spore release. However, the climatic conditions required for spore release are unclear. This is probably due to the difficulties involved in identifying and isolating the various types of spore (ascospores or macroconidia). Furthermore, the results obtained probably depend on the maturity of the spores: four periods of release over a period of 20 to 30 days have been observed.

However, it seems that macroconidia are dispersed by means of splashing during rainfall. In contrast, the release of ascospores is only triggered by rainfall (or high humidity), resulting in the rehydration of the perithecia after a dry period. The true release of the ascospores occurs several days later, depending on air humidity. Ascospore release, in *F. graminearum* in particular, appears to be periodic.

Although the dispersal pathways of the spores are known (splashing for the macroconidia and/or ascospores and wind dispersal for ascospores only) and long-distance dispersal appears to be possible, the maximum dispersal distance and the contaminating potential of splashing and wind dispersal have yet to be determined. Furthermore, local contamination seems to predominate in the epidemiological profile of the disease in that two neighbouring plots planted with the same variety may present very different levels of contamination.

The importance of local contamination raises the question of the hypothetical contamination of spikelets via the systemic route. This question remains unanswered because as far as we know no study has shown, microscopically, the systemic colonisation of the spikelets and the growth of the fungus in plant tissues (or the impossibility of such colonisation). Similarly, the existence, conditions of existence and potential importance of such contamination have not
been precisely determined. Conflicting results have been obtained and it is possible that this
route of contamination is possible only for a few plant genotypes, for a few fungal species or
strains, or for a few plant-fungus interactions. Indeed, interactions between plants and
pathogens have been observed during infection. We can therefore presume that the virulence
characteristics of the infecting fungal strain, the resistance of the plant and the virulence-
avirulence interaction between plants and pathogens determines the capacity of the plant to be
infected by mycelium and/or spores, the speed of infection (colonisation) of the ear by the
fungal mycelium and the amount of mycotoxins present in the grains at harvest. These
phenomena, like the production of inoculum, also depend on climate.

The choice of variety is a practical consideration that may affect plant infection: the date and
site of fungal penetration, the propagation rate and the intensity of the consequences of the
attack (severity of symptoms and/or amount of toxins). Indeed, the choice of variety
determines the major mechanisms of varietal resistance activated. However, it should be
stressed that although various types of resistance have been identified, they have not yet been
entirely elucidated. Furthermore, the types of varietal resistance have not yet been
characterised for most wheat varieties, which makes it difficult to compare varieties in a
particular area. Together with genetic characteristics, sowing date and climate determine the
date at which the crop is most susceptible to infection.

The extent of infection depends directly on the quality of inoculum, the primary source of
which is crop residues. In the epidemiological cycle of the disease, residues probably modify
the conservation and development of the fungus and spore dispersal, with effects depending
on the amount of residue present (which depends on soil tillage methods and the preceding
crop) and their nutritive value for the pathogen and for the plant (which also depends
essentially on the preceding crop). The high levels of nitrogen in maize residues may result in
a longer period of residue colonisation by *Fusarium*, strengthening pathogen populations, and
stimulating their development. If too many crop residues are present, spore dispersal by
spashing may be physically limited by obstacles. Conversely, if the fungus has access to too
few residues or residues insufficiently rich for it to complete its life cycle and to develop, then
the fungus is likely to produce fewer spores and to disperse less effectively. There is also an
interaction between these factors. Limited soil tillage methods increase the density of residues
on the surface of the soil and hence increase the quantity of inoculum available, particularly if
the preceding crop is a potential host of the fungus (e.g. maize or durum wheat). Mycelium
development and the production and dispersal of spores therefore depend on climate and irrigation.

Although there is broad agreement among scientists and advisers concerning the effects of soil tillage, irrigation and preceding crop on both disease severity and contamination by mycotoxins, the cause of these effects remains unclear. In addition, uncertainties remain concerning the survival of the fungus and its capacity to sporulate in soil, according to soil aeration, porosity and light penetration, all of which are affected by soil tillage. Similarly, the role of the compounds generated by the degradation of crop residues and the rate of residue degradation — which depend on soil tillage, the nutritive value of the preceding crop and the microclimate of the residues, which itself depends on irrigation and its relationship to mineral composition (nitrogen, silicon, depending on the preceding crop) — make it difficult to predict the quantitative effects of soil tillage, preceding crop and irrigation.

Other unanswered questions remain concerning the effect of residue degradation on competition between microorganisms and/or plant resistance (notably by means of differences in mineral nutrition).

The effect of mineral nutrition on disease severity and mycotoxin contamination, if indeed there is one, remains unclear. The mineral balance of the soil influences pathogen populations and should therefore have repercussions for the development of *Fusarium*. The composition of plants, in terms of nitrogen and silicon, for example, probably affects the growth and development of the fungus, although this has not been explicitly demonstrated in wheat for several reasons: equilibrium between the various elements, fertilisation date, link with *Fusarium* foot rot etc. A canopy with unlimited mineral nutrition develops well. This development slightly modifies the environmental conditions of the residues, which may in turn modify the rate of residue degradation. If this is indeed the case, then the capacity of the fungus to sporulate and of the spores to disperse may be affected.

The role played by canopy density, in terms of crop plant and weed densities, and in the development of epidemics of the disease remains to be determined. Weeds may carry *Fusarium*, but is their role limited to that of an intermediate host (during splashing) or do they act as a source of inoculum? If they serve as a source of inoculum, how many spores are produced on weeds? Weeds probably also play an indirect role, modifying mineral nutrition
and/or microclimate by increasing plant population density. The overall density of plants in
the canopy also depends on sowing density and the tillering capacity of the variety. Thus,
weeds may affect the number of spores produced, spore maturation and/or spore dispersal.
The uncertainties that remain probably arise at least in part from variations in the role played
by weeds according to the type or family of weeds concerned and competition with the crop:
date of emergence, density etc. (which also depends on climate). Although canopy density
(crop plus weeds) seems likely to play a role, we cannot be sure of the nature or magnitude of
this effect.

It also seems clear that competition between microorganisms on the leaves and ears is likely
to have an effect, but this is difficult to study in the field and necessarily depends on the
fungicides used and on canopy development.

Five types of varietal resistance have been identified to date. They affect the penetration of
the fungus into the plant (type I), infection kinetics (type II), the expression of the infection
(symptoms; type III), the consequences of infection (quantity of mycotoxins produced, yield;
type IV) and grain infection (type V). This complexity of plant resistance complicates any
study of several varieties. In such studies, caution is required when interpreting results
analysing the relationships between cultural practices and symptoms, between symptoms and
the quantities of mycotoxins produced and between cultural practices and the mycotoxins
produced.

In conclusion, the relationship between disease symptoms and mycotoxin contamination is of
key importance. The conditions in which this relationship is purely qualitative and those in
which it also has a quantitative element are unclear. If we are to propose effective methods for
the prevention of grain contamination by toxins, we must improve our understanding of this
relationship, especially as concerns the role of the cropping system. Investigation of the
effects of crop management and crop rotation on the profile of *Fusarium* species and strains
obtained within a cereal field, and the conditions in which the potential for toxin production is
expressed in agricultural environments, should be a research priority.

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Figure 1: Symptoms of *Fusarium* head blight on the external surface of wheat ear glumes.
Table 1: Data on environmental conditions for the production of macroconidia and ascospores in the laboratory, according to species. Two natural populations of *F. graminearum* can be identified: Group 1 (or *F. pseudograminearum*), normally associated with the diseased crowns of host plants, which do not form perithecia in culture and form such structures only rarely in nature, and Group 2, associated with diseased of aerial plant parts, which do form perithecia- [2,223,224].
<table>
<thead>
<tr>
<th>Stage</th>
<th>Species</th>
<th>Determinant factors</th>
<th>Range</th>
<th>Optimal value</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mycelial growth</td>
<td><em>F. graminearum</em> group 2 (aerial)</td>
<td>Pressure, Humidity</td>
<td>&gt; -9 bars, &gt; 94%RH</td>
<td></td>
<td>[75] (on maize grains)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pressure</td>
<td>&gt; -12 bars, -2 bars</td>
<td></td>
<td>[225] (on agar)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Temperature</td>
<td>4 to 32°C, 28°C</td>
<td></td>
<td>[47]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Temperature</td>
<td>20 to 30°C, -10 to –28 bars</td>
<td></td>
<td>[23] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Accelerated by alternate periods of rainfall and drought before ear emergence</td>
<td>[59]</td>
</tr>
<tr>
<td></td>
<td><em>F. culmorum</em></td>
<td>Pressure</td>
<td>20 to 30°C, -8 to –14 bars</td>
<td></td>
<td>[23] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pressure</td>
<td>At –8 bars, T. optim 20°C, At –14 bars T. optim 30°C, At –28 bars, T. optim 38°C</td>
<td></td>
<td>[16]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Temperature</td>
<td>5 to 35°C, 29°C</td>
<td></td>
<td>[2,16]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pressure</td>
<td>&lt; -50 bars, poor from –5 bars, -1.5 bars</td>
<td></td>
<td>[97] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Temperature</td>
<td>0 to 28-32°C, 18-20°C</td>
<td></td>
<td>[11, 16]</td>
</tr>
<tr>
<td>Initiation of perithecia</td>
<td><em>F. graminearum</em> group 2 (aerial)</td>
<td>Light</td>
<td>Requires low intensity of UV light (300-320&lt;390nm)</td>
<td></td>
<td>[83], with conflicting results obtained by [35].</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Depends on the strain</td>
<td></td>
<td>[35,71]</td>
</tr>
<tr>
<td></td>
<td><em>F. roseum</em></td>
<td>Light</td>
<td>Requires low intensity of UV light (300-320&lt;390nm)</td>
<td></td>
<td>[83], with conflicting results obtained by [35].</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Depends on light</td>
<td></td>
<td>[59]</td>
</tr>
<tr>
<td>Production of perithecia</td>
<td><em>F. graminearum</em> group 2 (aerial)</td>
<td>Light</td>
<td>Requires low intensity of UV light (300-320&lt;390nm)</td>
<td></td>
<td>[83], with conflicting results obtained by [35].</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Depends on light</td>
<td></td>
<td>[59]</td>
</tr>
<tr>
<td></td>
<td><em>M. nivale</em></td>
<td>Temperature</td>
<td>&lt; 16-18 °C, 6 à 8 °C</td>
<td></td>
<td>[64]</td>
</tr>
<tr>
<td>Production of spores</td>
<td><em>Fusarium</em></td>
<td>Temperature</td>
<td>Around 10°C, 15°C</td>
<td></td>
<td>[64]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Humidity</td>
<td>Around 80% humidity</td>
<td></td>
<td>[64]</td>
</tr>
<tr>
<td>Production of macroconidia</td>
<td><em>F. graminearum</em> group 2 (aerial)</td>
<td>Temperature</td>
<td>28 to 32°C, None if T°&lt;16°C or T°&gt;36°C</td>
<td></td>
<td>[83]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pressure</td>
<td>max &lt; –50 bars, -1.4 to –3 bars</td>
<td></td>
<td>[97] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Temperature</td>
<td>16-36°C, 28-32°C</td>
<td></td>
<td>[47]</td>
</tr>
<tr>
<td></td>
<td><em>F. culmorum</em></td>
<td>Temperature</td>
<td>20 to 30°C, -10 to –28 bars</td>
<td></td>
<td>[23] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pressure</td>
<td>Max between -15 and –60 bars, -15 bars</td>
<td></td>
<td>[97] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td><em>F. graminearum</em> group 1 (soil)</td>
<td>Pressure</td>
<td>Max between -15 and –60 bars, -15 bars</td>
<td></td>
<td>[97] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td><em>F. avenaceum</em></td>
<td>Pressure</td>
<td>Max between -15 and –60 bars, -15 bars</td>
<td></td>
<td>[97] (on osmotically adjusted agar)</td>
</tr>
<tr>
<td></td>
<td><em>F. roseum</em></td>
<td>Light</td>
<td>UV light required</td>
<td></td>
<td>[16]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Humidity</td>
<td>RH of about 100% Favoured by water stress</td>
<td></td>
<td>[16]</td>
</tr>
<tr>
<td></td>
<td><em>M. nivale</em></td>
<td>Temperature</td>
<td>1 to 16 °C, 6 to 8 °C</td>
<td></td>
<td>[16]</td>
</tr>
</tbody>
</table>
Table 2: Spectrum of infection of weeds by *Fusarium* species. M: Monocotyledonous, g: grasses, D: Dicotyledonous.

<table>
<thead>
<tr>
<th>Weed host Family</th>
<th>Latin name</th>
<th>Common name</th>
<th><em>Fusarium</em> Species</th>
<th>Site and form</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poaceae Mg</td>
<td><em>Agropyron</em></td>
<td>Wheatgrass</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. graminearum</em></td>
<td>Not indicated</td>
<td>[85]</td>
</tr>
<tr>
<td>Liliaceae M</td>
<td><em>Allium</em></td>
<td>Garlic</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. graminearum</em></td>
<td>Not indicated</td>
<td>[64]</td>
</tr>
<tr>
<td>Poaceae Mg</td>
<td><em>Alopecurus myosuroides</em> Huds.</td>
<td>Blackgrass</td>
<td><em>F. roseum</em></td>
<td>In the seed</td>
<td>[48]</td>
</tr>
<tr>
<td>Poaceae Mg</td>
<td><em>Avena fatua</em> L.</td>
<td>Wild oats</td>
<td><em>F. roseum</em></td>
<td>Lesion, spores on leaves and seeds</td>
<td>[48]</td>
</tr>
<tr>
<td>Chenopodiaceae D</td>
<td><em>Beta vulgaris</em></td>
<td>Wild beet</td>
<td><em>F. culmorum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Cruciferae D</td>
<td><em>Capsella bursapastoris</em> L.</td>
<td>Shepherd's purse</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. poae</em>, <em>F. sambucinum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Compositae D</td>
<td><em>Cirsium arvense</em> L.</td>
<td>Thistle</td>
<td><em>F. avenaceum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Poaceae Mg</td>
<td><em>Dactylis</em> L.</td>
<td>Cocksfoot</td>
<td><em>F. roseum</em></td>
<td>Lesion on straw, stem base, increase with N</td>
<td>[14, 48]</td>
</tr>
<tr>
<td>Poaceae Mg</td>
<td><em>Echinochloa crus-galli</em> L.</td>
<td>Barnyardgrass</td>
<td>Not indicated</td>
<td>Not indicated</td>
<td>[85]</td>
</tr>
<tr>
<td>Poaceae Mg</td>
<td><em>Festuca</em></td>
<td>Fescue</td>
<td><em>F. roseum</em></td>
<td>Stem base, increase with N</td>
<td>[14]</td>
</tr>
<tr>
<td>Rubiaceae D</td>
<td><em>Galium aparine</em> L.</td>
<td>Cleavers</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. poae</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Poaceae Mg</td>
<td><em>Lolium multiflorum</em> L.</td>
<td>Ryegrass</td>
<td><em>F. roseum</em></td>
<td>Lesion on straw, stem base, increase with N</td>
<td>[14, 48]</td>
</tr>
<tr>
<td>Compositae D</td>
<td><em>Matricaria spp.</em></td>
<td>Mayweed</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. graminearum</em>, <em>F. poae</em>, <em>F. sambucinum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Ranunculaceae D</td>
<td><em>Ranunculus acris</em></td>
<td>Common buttercup</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. poae</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Ranunculaceae D</td>
<td><em>Ranunculus repens</em> L.</td>
<td>Creeping buttercup</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Polygonaceae D</td>
<td><em>Rumex obtusifolius</em></td>
<td>Dock</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Compositae D</td>
<td><em>Senecio vulgaris</em> L.</td>
<td>Groundsel</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Caryophyllaceae D</td>
<td><em>Spergula arvensis</em> L.</td>
<td>Corn spurrey</td>
<td><em>F. sambucinum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Caryophyllaceae D</td>
<td><em>Stellaria media</em> L.</td>
<td>Chickweed</td>
<td><em>F. avenaceum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Fabaceae D</td>
<td><em>Trifolium</em></td>
<td>Clover/trefoil</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. graminearum</em></td>
<td>Not indicated</td>
<td>[64]</td>
</tr>
<tr>
<td>Urticaceae D</td>
<td><em>Urtica dioica</em> L.</td>
<td>Nettle</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. sambucinum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Scrophylariaceae D</td>
<td><em>Veronica persica</em></td>
<td>Speedwell</td>
<td><em>F. graminearum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Violaceae D</td>
<td><em>Viola arvensis</em> Murray</td>
<td>Field violet</td>
<td><em>F. avenaceum</em>, <em>F. poae</em>, <em>F. graminearum</em>, <em>F. sambucinum</em></td>
<td>Stem base</td>
<td>[62]</td>
</tr>
<tr>
<td>Cryophyllaceae D</td>
<td><em>Dianthus</em></td>
<td>Carnation</td>
<td><em>F. avenaceum</em>, <em>F. culmorum</em>, <em>F. poae</em>, <em>F. graminearum</em></td>
<td>Not indicated</td>
<td>[64]</td>
</tr>
</tbody>
</table>
Figure 2: Chemical structures of trichotecenes, fumonisin B1 and zearalenone.

**Trichotecenes**

Type A trichotecenes:  
- T-2 (R1 = OAc)  
- HT-2 (R1 = OH)

Type B trichotecenes:  
- DON (R1 = OH, R2 = H, R3 = OH, R4 = OH)  
- NIV (R1 = OH, R2 = OH, R3 = OH, R4 = OH)

**Fumonisin B1**

**Zearalenone**