Mini-Review article

**Cylindrocarpon destructans/Ilyonectria radicicola-species complex: Causative agent of ginseng root-rot disease and rusty symptoms**

Mohamed El-Agamy Farh¹, Yeon-Ju Kim²,*, Yu-Jin Kim², Deok-Chun Yang¹,²,*

¹ Graduate School of Biotechnology and Ginseng Bank, College of Life Science, Kyung Hee University, Yongin, Republic of Korea
² Department of Oriental Medicinal Biotechnology, College of Life Sciences, Kyung Hee University, Republic of Korea

**A R T I C L E   I N F O**

Article history:
Received 8 September 2016
Received in Revised form
11 November 2016
Accepted 2 January 2017
Available online 16 January 2017

Keywords:
*Cylindrocarpon destructans*
*I. radicicola* species complex
Panax ginseng
root-rot
rusty root

**A B S T R A C T**

*Cylindrocarpon destructans/Ilyonectria radicicola* is thought to cause both rusty symptom and root-rot disease of American and Korean ginseng. Root-rot disease poses a more serious threat to ginseng roots than rusty symptoms, which we argue result from the plant defense response to pathogen attack. Therefore, strains causing rotten root are characterized as more aggressive than strains causing rusty symptoms. In this review, we state 1- the molecular evidence indicating that the root-rot causing strains are genetically distinct considering them as a separate species of *Ilyonectria*, namely *I. mors-panacis* and 2- the physiological and biochemical differences between the weakly and highly aggressive species as well as those between rusty and rotten ginseng plants. Eventually, we postulated that rusty symptom occurs on ginseng roots due to incompatible interactions with the weakly aggressive species of *Ilyonectria*, by the established iron-phenolic compound complexes while root-rot is developed by *I. mors-panacis* infection due to the production of high quantities of hydrolytic and oxidative fungal enzymes which destroy the plant defensive barriers, in parallel with the pathogen growth stimulation by utilizing the available iron. Furthermore, we highlight future areas for study that will help elucidate the complete mechanism of root-rot disease development.

© 2017 The Korean Society of Ginseng, Published by Elsevier Korea LLC. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

1. **Introduction**

The genus *Panax* is a member of the Araliaceae family. The name *Panax*, given by Carl A. Meyer, originates from the Greek word meaning “cure-all” and includes members used for many decades as oriental medicinal plants [1,2]. There are 11 accepted species in the genus *Panax* (http://www.theplantlist.org/browse/A/Araliaceae/Panax/). Six of these species are well-known: *P. quinquefolius* and *P. trifolius*, which are native to North America [3,4], and *P. ginseng*, *P. notoginseng*, *P. vietnamensis* and *P. japonicus* which are located in Asia. *P. quinquefolius*, *P. ginseng*, and *P. notoginseng* are considered important cash crops due to their pharmaceutical properties [5,6]. These pharmaceutical properties are due to the major components of the ginseng root, dammarane-type saponins, which are referred to as ginsenosides [1,2,7].

Ginseng plants are slow-growing perennial herbs that require several continuous growing seasons for optimal growth and ginsenoside accumulation. Four- to six-year-old ginseng roots have the best shape, growth, and ginsenoside content [8]. Accordingly, optimal growth is required over a long cultivation period before harvesting.

During the six years of cultivation, ginseng plants can suffer from several diseases that lead to dramatic crop losses and reduction of root quality [9–11]. Diseases can be divided into two types: foliar-borne diseases and soil-borne diseases. Examples of foliar disease are blight, anthracnose, and mildew. Blight can occur in different forms. The first form appears on the leaves during the initial stage as circular, water-soaked spots. Over time, the spots dry and turn brown with a targeted-board appearance and clear dark brown margin. This type of blight is caused by *Alternaria panax* and accordingly is called *Alternaria* blight [8,12–14]. The second form appears mostly at the leaf tip and expands backward through the midrib, resulting in a characteristic V-shape infection. The lesions have a targeted-board appearance like *Alternaria* infection;
however, at the late stage of infection, the lesions cover the leaf surface, the color of the leaves becomes reddish to brown, and the pathogen grows densely and sporulates, giving the infected leaves a characteristic gray color. This blight disease can be differentiated from *Alternaria* blight by its ability to infect flowers and fruits. This blight is caused by *Botrytis cinerea* and is accordingly called *Botrytis* blight [12,15]. Anthracnose disease is found on leaves exposed to direct sunlight. The symptoms of this disease are similar to those of *Alternaria* blight; however, anthracnose disease is characterized by a felt-like appearance. The causal agent of anthracnose disease has been reported to be *Colletotrichum panaciciola* [12,16,17]. Mildew disease had been considered as a type of blight; however, the name was changed to refer to symptoms completely different from those of *Alternaria* and *Botrytis* blights. At the early stage of disease, the blights appear similar to *Alternaria* and *Botrytis* blights, but later, the blight turns to dark green with a whitish center. In addition, mildew blights lack the yellowish-brown margins seen in *Alternaria* blights or a grayish surface as seen in *Botrytis* blights. Mildew disease is induced by *Phytophthora cactorum* [12,18–20].

Soil-borne diseases include damping-off, rusty root, and root-rot disease. Damped-off ginseng plants usually have intact and erect stems with drooping leaves. Leaves finally turn yellowish, the stem collapses, and the plant dies. Damping-off disease has been reported to be induced by *Pythium* spp. [21]. Rusty root disease is usually characterized by small or quite large reddish brown areas at the crown of the tap root that can be easily scraped off to show the inner white healthy tissue, while rotten root disease is characterized by the complete absence or only partial presence of hollow roots with dark brown discoloration. Interestingly, rusty root disease and root-rot disease are both caused by *Cylindrocarpon destructans* var. *destructor* [21–23]. Although foliar diseases cause plant losses, the root mostly remains intact and is not physically harmed. Sometimes, limited growth or lowered production of bioactive metabolites is observed because of defoliation and reduction in leaf area, as in the case of *A. panax* blight [8], but overall, the reduction in yield is minimal. Root diseases, in contrast, pose the greatest threat to ginseng cultivation and crop yield because the infection directly harms the roots, leading to changes in root shape and reduction in root quality, in addition to physiological changes [24,25]. Damping-off disease threatens the roots of ginseng crop at the seedling stage; however, careful handling and treatment during ginseng cultivation at this stage can prevent this disease from causing damage, as the causal agent cannot invade older roots [21]. Root-rot and rusty root diseases are the most dangerous diseases for ginseng crops as they cause a great decrease in yield and damage root shape and quality in plants of all ages [25]. As explained earlier, these two diseases have been attributed to infection by *Cylindrocarpon destructans var. destructans* [22,25]. Previous studies have reported that this pathogen is genetically diverse and can be divided into several species [26]. In this review, we detail the main morphological differences between these two diseases and environmental factors leading to their development. Furthermore, we detail the re-classification of this pathogen into several species, and we describe, for the first time, how this correlates with the morphological and biochemical characteristics of root-rot and rusty root-causing strains. In addition, we discuss the mechanism of root-rot and rusty root development. Finally, we highlight points that need to be clarified in the future for a comprehensive understanding of how root-rot is initiated and measures that can be taken to combat this disease.

2. Description of ginseng root-rot and rusty root diseases

Root-rot is a soil-borne disease that causes huge loss of ginseng crops. The disease is detected at all stages of plant growth. In addition, the causal agent has the ability to overwinter in the soil in a resting stage, so that newly planted crops and existing crops can be re-invaded each season [12,22,23,25]. Therefore, it is considered the most serious disease in most ginseng cultivation areas. Root-rot disease in American ginseng was first described by Zinsmeister [27]. Similar symptoms were then observed on Korean ginseng, and the same causative pathogen was discovered by Chung [28]. The disease usually appears as a dark brown discolored area on the tip of the tap root and extends over time upward to the crown. Sometimes, rotting can affect any part of the root. Due to invasion, the outer surface of the plants is damaged, while the inner part totally disintegrates, leaving a hollowed-out root in the soil (Fig. 1). At the late stage of root-rot disease, the leaves turn yellow, wilt, and

![Fig. 1. A schematic showing the morphological development of root-rot and rusty root symptoms.](image-url)
finally the end of the stem get easily separated from the crown.
Because of these characteristic symptoms, this kind of rot disease is
called disappearing root-rot [12,25]. The causal agent of such
symptoms have been reported to be, as previously named, C. destructans var. destructans [21–23]. Fusarium species exist in
association with root-rot symptoms; however, there is doubt that
they are the primary causal agents of rooting disorder [25] as
artificial infection of ginseng seedlings by Fusarium species causes
root-rot [29], and these species have not been reported to cause
rotting symptoms in older roots. Another uncommon root disease
that has been detected on ginseng root differs from Cylindrocarpon
root-rot by the absence of brown discoloration and the presence of
soft, water-soaked rot in the cortex. This type of rotting is caused by
Phytophthora cactorum [12].

Rusty root or rusted root is another common disease that affects
ginseng roots at all stages of growth. The disease has been reported
in North American countries, e.g., Canada, as well as in Asian
countries, e.g., South Korea and China [12,24]. Infected roots have
reddish scabs covering the entire root or part of the root. These
scabs do not cause plant death; however, they can reduce root
quality and price by up to 40%. Symptoms usually start at the crown
of the tap root as small or large, raised reddish-brown areas and can
sometimes be present on other parts of the root (Fig. 1). The
symptoms are usually superficial, and can be easily scraped off to
show the inner, white healthy tissues [12,24,25]. Rusty root disease
had been observed to be associated with C. destructans var.
destructans in particular, weak strains of C. destructans var.
destructan are unable to cause intensive root rotting [22,25].

3. Causal agent

3.1. Taxonomy of the pathogen; from single species to species
complex

The genus Cylindrocarpon was originally proposed by Wollenweber [30] as the anamorph of the teleomorphic state of the Nectria
section Willkommiiotes Wollenw. and was characterized by the
inability to produce chlamydospores. A few years later, the generic
name was expanded to include species able to produce chla-
mydospores, including C. destructans. Then, Booth [31] divided the
genus into four different groups based on the presence or absence
of microconidia and chlamydospores: C. magnusianum (Sacc.)
Wollenw.; C. cylindroides Wollenw., the type species of the genus
Cylindrocarpon; the group of Cylindrocarpon species considered the
anamorph of Nectria mammoides; and C. destructans, the anamorph of
N. radicicola. C. destructans is characterized by the presence of
both microconidia and chlamydospores. Later, morphological
variation among C. destructans strains was observed, and their
teleomorphs were divided into varieties accordingly. Those
infecting ginseng plants were given the name C. destructans var.
destructans, and the teleomorph is named N. radicicola var. radici-
cola [32]. By this time, all Nectria species with the anamorph
Cylindrocarpon were included in the genus Neonectria [33–35].

Based on the morphological characteristics of the teleomorph of
the genus Cylindrocarpon, they were divided into five informal
groups: (1) Neonectria coccinea/galligena-group (Neonectria sensu
stricto) [36]; (2) N. mammoides-group [36]; (3) N. rugulosa-group
[37]; (4) N. radicicola-group [36]; and (5) N. veillottleina-group [38].

This classification was revised based on analysis of mitochondrial
DNA data into three different clades; the C. destructans/N. radicicola
group formed clade III [34]. The morphological and molecular
variation found among Cylindrocarpon and its teleomorph, Neo-
ectria, suggested the presence of several genera within the genus
Cylindrocarpon/Neonectria. Multilocus analysis [e-actin (ACT), in-
ternal transcribed spacer (ITS), large subunit (LSU), RNA
polymerase II subunit 1 (rpb1), translation elongation factor 1–z
(TEF-1–z), β-tubulin (TUB)] revealed the presence of five different
groups that overlapped with both Booth [31] and Mantiri [34]
classifications. The phylogenetic analysis correlated with observed morphological differences, particularly macroconidial
septation and perithecia anatomy. Therefore, each group was
considered a distinct genus, and the N. radicicola group was given
the generic name Ilyonectria. Accordingly, N. radicicola var. radici-
cola (the teleomorph of C. destructans var. destructans) was named
I. radicicola [39]. The description of I. radicicola in this study is based on
the amended description of Booth [31,40] and Samuels and
Brayford [32]; perithecia usually solitary with papillate, conical
shape apex, smooth to slightly roughed surface containing 1–
septate smooth ascospores; anamorph produces straight cylin-
drical macroconidia with rounded ends and conspicuous hilum,
elipsoidal microconidia with or without conspicuous hilum;
chlamydospores intercalary, single or in chains, and becoming
brown after aging. A few months later, it was proposed that the
anamorph of I. radicicola is a species complex; multigene analysis
[TUB, histone H3 (HIS), TEF-1–z, and nuclear ribosomal RNA-
internal transcribed spacer (nrRNA-ITS)] of I. radicicola ana-
mosraphs isolated from many hosts proved that members of that
genus are not monophyletic. Rather, they clustered in many groups,
each of which was considered a separate species. In this study, both
teleomorphic and anamorphic states are referred to by a single
generic name, as for the other pleomorphic fungi [41–45]. Of 68
isolates obtained from many hosts, 21 isolates obtained from
ginseng plants were resolved into four different Ilyonectria species:
I. mors-panacis, I. robusta, I. panacis, and I. crassa [26]. The taxo-
nomic position of ginseng plant-infecting Cylindrocarpon within
each classification study from 1966 to the present is illustrated in
Fig. 2. Despite the high morphological similarity among the four
resolved Ilyonectria species, some differences have been observed,
particularly in the morphology and size of the conidia, in addition to
colony diameter at 25°C (Table 1).

3.2. Correlations between morphological, biochemical, and genetic
diversity and pathogenicity

Before the Cabral [26] classification, many differences in
morphology, optimal temperature for growth, enzymatic activity,
and genetic profiles were detected among strong and weak strains
and were assumed to be linked to the observed differences in
pathogenicity. Rahman and Punja [25] observed that highly
and weakly aggressive strains were morphologically different;
the highly aggressive strain had a dark to rust brown-colored ap-
pearance after two weeks of growth on PDA media, while the weakly
aggressive strain had a beige to light brown-colored appearance.
Additionally, the optimal temperature for growth differed between
the strains; they observed that the optimal radial growth of highly
aggressive and weakly aggressive strains on PDA media plates
occurred at 18 and 21°C, respectively. Production of enzymes to
dergrade plant cell wall materials (e.g., pectinase and cellulase) and
oxidative enzymes to detoxify the accumulated polyphenolic
compounds by highly and weakly aggressive was also character-
istic; Rahman and Punja [25] reported that virulent strains pro-
duced pectinase and polyphenol oxidase in greater quantities than
avirulent strains. In another study, it was proposed that root-rot-
casing strains are able to produce cellulose in high quantities,
but the ability of weakly aggressive strains to produce cellulose has
not been explored [46]. The morphological and biochemical dif-
fferences between highly and weakly aggressive strains suggested
that the highly aggressive strains are genetically distinct. Combined
phylogenetic analysis of both ITS and TUB from weak and aggres-
sive strains revealed that the aggressive strains were
and were given the name aggressive strains were therefore considered to be a forma specialis similar to those originally infect American ginseng in Canada. The American ginseng. In addition, they were found to be genetically ginseng in Korea and Japan were recorded as aggressive strains on type factor appear to play a role in the genetic distinctiveness of the phylogenetically distinct. Neither geographical location nor host derived aggressive species were found to be distinct. On the other hand, ginseng-derived weakly aggressive strains were found to be genetically similar to the strains derived from different hosts. For example, I. robusta derived also from the roots of Loroglossum hir-cinum, Tilia petiolaris, Quercus robur, Prunus cerasus, Vitis vinifera, and Thymus spp. I. crassa has also been observed on the roots of Narcissus spp. and bulbs of Lilium spp. [26]. The identity of the original host of I. panacis is unclear, as only one ginseng-derived strain has been investigated in detail. These findings suggest that American and Korean ginseng are not susceptible hosts to I. robusta and I. crassa. In addition, there may be a host-specific interaction between I. mors-panacis and its sole host, ginseng.

4. Environmental factors involved in disease development

Beside the main biotic factor, the pathogen, the density of root-rot disease in ginseng crops is influenced by many abiotic factors that can either favor pathogen growth or increase plant susceptibility to pathogen infection. For example, if wounds made by soil insects or cultivation tools exist on the root surface, they may play a role in root-rot development as weak points through which the pathogen can invade the root aggressively; wounded roots have been found to be more susceptible to infection than unwounded ones [25]. Soil pH also has an effect on disease intensity; generally, the pH of soils with good ginseng root yields is weakly acidic (5.5-6) [48,49], while more acidic soil (pH < 5) has been observed to be associated with disease abundance. This is consistent with the observation that root-rot-causing I. radicicola-species complex of conidiophores

**Table 1** Summary of the morphological differences between Ilyonectria radicicola-species complex infecting American and Korean ginseng [26]

| Type of conidia | characteristic | I. crassa | I. robusta | I. panacis | I. mors-panacis |
|-----------------|----------------|-----------|------------|------------|----------------|
| Macroconidia    | Origin of production | Produced on simple complex of conidiophores | Produced on simple conidiophore | Produced on simple complex of conidiophores | Produced only on the simple conidiophores |
| Shape of ends   | Both ends are rounded, but sometime narrowing at the tip | Hilum exist | No hilum exist | Hilum exist | No hilum exist |
| Hilum existence | 25.6 × 1.5 | 23.7 × 1.5 | 24.8 × 1.5 | 29.9 × 1.5 | 34.4 × 1.5 |
| Mean size of (μm) | 2- to 3-septate | 27.2 × 1.5 | 28.7 × 1.5 | 41.0 × 1.5 |
| Microconidia    | Hilum exist | No hilum exist | Hilum exist | Hilum exist | Hilum exist |
| Origin of production | Produced only on the simple conidiophores | Both ends are rounded | Both ends are rounded |
| Shape of ends | Both ends are rounded | Both ends are rounded | Both ends are rounded |
| Hilum existence | No hilum exist | No hilum exist | No hilum exist | No hilum exist | No hilum exist |
| Mean size of (μm) | 4.9 × 1.5 | 5.2 × 1.5 | 5.6 × 1.5 | 5.1 × 1.5 | 5.7 × 1.5 |
| Growth at temperature | 19-34 | 35-48 | 15 | 31-40 |

**Fig. 2.** Schematic overview of the taxonomic position of ginseng plant-infecting Cylindrocarpon species within each classification study from Booth [32] until Cabral [26]. Blue-colored squares refer to the group of Cylindrocarpon/Ilyonectria in which those infecting ginseng plants are included. Dotted circle shows the Ilyonectria species, I. mors-panacis that has been reported only in ginseng plants.
example, the foliar application of iron [50] and absence of calcium [51] in ginseng roots that have been observed to occur in the temperature range of 13-23°C. Another factor that determines the severity of root-rot disease is the age of the root; although the disease can be found in all ages of ginseng plant, young roots, in particular two-year-old-roots, appear to be the most susceptible [25], possibly because the root cell wall is still immature and not yet well developed. Temperature also plays a critical role in determining the intensity of root-rot disease; rotting symptoms were observed to occur in the temperature range of 13-23°C, while no symptoms were observed at 28°C [52].

Environmental factors that influence rusty root development are similar to those that influence root-rot development, in particular, the presence of metal ions in the ginseng soil; high content of certain cations (e.g., iron, aluminum, and silicon) has been noted in rusty root lesions versus healthy tissues, which conversely have high calcium and potassium levels [24,53]. It was deduced that such metal variation, in combination with pathogen infection, may be responsible for the appearance of rust.

5. Mechanism of infection and disease development

As mentioned earlier, rusty root disease causes less damage to ginseng roots than root–rot [25]. Rusty root symptoms have also been reported to be caused by other fungi, e.g., Fusarium spp. [54], Rhoxercosporidium carota [55], Pseudomonas marginalis, Micro-bacterium oxydans, Lysobacter, Gommonos, Rhizobium leguminosarum, Pseudomonas aeruginosa, and Agrobacterium tumefaciens [56]. Therefore, rusty-colored lesions on the root surface may represent a plant’s defensive response to invasion by microorganisms that are unable to cause root rot. As rusty symptoms are economically less important than root-rot disease, it is important to elucidate the interaction between ginseng roots and the two groups of Ilyonectria that cause rusty root and root-rot symptoms to determine how to manage root-rot-causing pathogens and limit disease. In rusty root, the levels of phenolic compounds and iron are much higher in the rusty tissues than adjacent healthy tissues [24,53]. In addition, production of other defense-related enzymes, e.g., phenylalanine ammonia-lyase, polyphenoloxidase, and peroxidase, is stimulated in rusty tissue. The formation of iron-phenolic compound complexes is thought to be the reason for the rust color formation [24]. Rust-causing Ilyonectria species produce low levels of plant cell wall hydrolytic enzymes and phenolic-detoxifying enzymes [25]. There is little information about physiological and biochemical changes in root-rot infected roots, but the pathogen can produce large amounts of hydrolytic enzymes and is able to detoxify phenolic compounds [25]. In addition, iron has been found to be essential for root-rot development and pathogen growth [50].

These findings indicate that rusty symptoms are not actually a sign of disease but the result of incompatible interactions between ginseng root and microorganisms, e.g., L. robusta, L. crassa, and L. panacis and others; the ability of these organisms to degrade the plant cell wall is very low; therefore, when they attempt to invade the root surface, they are not very successful, and the process is slow. The plant responds by producing defense-related enzymes, e.g., phenylalanine ammonia-lyase, polyphenoloxidase, and peroxidase, and secondary metabolites such as phenolic compounds. As these fungi do not have sufficient ability to detoxify these phenolic compounds, invasion is retarded. We argue that root-rot occurs due to compatible interactions between ginseng root and the aggressive Ilyonectria species L. mor-panacis; when the pathogen spore or mycelium attaches to the root surface, it rapidly produces high quantities of hydrolytic enzymes such as cellulase and pectinase, allowing rapid invasion of the epidermal layer and fast extension of the inoculum to the cortical and inner tissues. As the plant starts to respond to the invasion by producing phenolic compounds, the pathogen responds by producing enzymes that can break down phenolic compounds such as polyphenoloxidases. In parallel, the pathogen sequesters iron from the plant to support its growth using siderophores. As the cell wall components degrade, the plant defense response declines, and the growth of the pathogen is supported by the plant and rhizosphere iron; eventually, all barriers to defeat the invading pathogen are overcome, and rotting symptoms are established (Fig. 3).

6. Summary and future directions

**Cylindrocarpon destructans** var. destructans is associated with both rusty and rotting symptoms in ginseng roots. Rusty symptoms do not appear to reflect disease, but to result from defense of ginseng roots to incompatible interactions with soil microorgan-isms [24]. Therefore, C. destructans var. destructans strains that cause rusty symptoms are unable to cause root-rot disease [22,25]. This raises several questions. Why are there two different kinds of symptoms caused by the same fungus? Are the fungus strains genetically similar or distinct? If distinct, what is the extent of the genetic differences? Are they different genotypes of the same species or different species? Finding the answers to these questions may explain how aggressive strains cause severe rotting and suggest suitable strategies to limit disease development. These questions started to be investigated when it was proposed that strains causing ginseng root-rot have a particular genotype. These strains were accordingly named C. destructans f. sp. panacis [47]. Nevertheless, the teleomorph of genus Cylindrocarpon was segregated into four different groups. Each group was considered to be a different genus, and C. destructans var. destructans teleomorph was renamed Ilyonectria radicicola [39]. Shortly thereafter, it was reported that the anamorphic state of I. radicicola is genetically polyphyletic. In addition, the genetic differences correlated with morphological variation, which provided sufficient evidence to consider each phylogenetic group to be a different species of Ilyonectria. Strains infecting ginseng plants were segregated into four different species: L. crassa, L. robusta, L. panacis, and L. mor-panacis. Interestingly, strains that cause severe root-rot diseases were clustered in one group: the L. mor-panacis group [26]. However, this correlation did not resolve the pathogenicity of other species complexes of Ilyonectria on their hosts. For example, L. liriodendri strains infecting grape tree have been reported to be a species complex, but their virulence does not correlate with genetic differences [57]. Other than genetic diversity, several pathological criteria have been used to distinguish the aggressive Ilyonectria species, L. mor-panacis, from others, e.g., the production of high quantities of hydrolytic and oxidative enzymes to rapidly degrade plant cell wall components and detoxify phenolic compounds produced by plant roots upon invasion. However, additional studies need to be performed to fully clarify the mechanism of L. mor-panacis infection. For example, the role of the saponins in disease development should be investigated; generally, saponins have antifungal activity and accordingly protect plants from soil-borne fungi [58–62]. Many studies have demonstrated the ability of some plant pathogens to tolerate the lethal activity of saponins, possibly through the production of saponin hydrolytic enzymes [60,63]. No studies have investigated if ginseng saponins play a role in plant protection. Only a few studies have investigated the role of ginsenosides in the pathogenicity of Pythium irregular, which causes damping-off of ginseng seedlings [62,64–66]; these studies concluded that the pathogenicity of this oomycetous pathogen depends on ginsenoside degradation [60]. Therefore, it is unknown how L. mor-panacis interacts with ginsenosides to negate their
antifungal properties. Does it degrade the ginsenoside or has another mechanism? Furthermore, the role of reactive oxygen species (ROS) during disease progression of *I. mors-panacis* needs to be investigated; in general, ROS are produced in plant cells when the plant is exposed to abiotic or biotic stresses [67,68] and act either as signaling molecules or cause oxidative damage, depending on their concentration [69]. When some pathogenic fungi invade plants, they cause fluctuations in ROS production. For example, the necrotrophic fungus *Botrytis cinerea* performs necrotrophic invasion through rapid induction of ROS while being resistant to the oxidative effect of these ROS, leading to the death of plant tissues [70]. In contrast, the semi-biotrophic fungus *Ustilago maydis* deals with ROS by inhibiting the activity of the peroxidase enzyme, which leads to suppression of ROS production and accordingly a decrease in defense-related signaling [71]. It must therefore be determined how *I. mors-panacis* copes with ROS. Investigating these points will increase our understanding of the mechanism of root-rot development and provide information that can potentially be exploited to manage and limit disease occurrence.

7. Conclusion

In this review, we conclude the genetic diversity among *Ilyonectria radicicola*-species complex infecting ginseng plants which state root-rot causing strains as a separate species named *I. mors-panacis* from others associated with rusty symptoms which named *I. crassa*, *I. robusta*, *I. panacis*. Furthermore, we figure out the physiological and biochemical differences between highly and weakly aggressive species as well as those between the rusty and rotten roots to elucidate the mechanism of the two diseases symptoms developed by weakly and highly aggressive species infection, respectively. Accordingly, we postulated that the rusty symptoms is developed during the incompatible interaction between ginseng roots and weakly aggressive microorganisms, including the species of *Ilyonectria* other than *I. mors-panacis*, possibly by the established iron-phenolic compound complexes while rotting symptom is developed during the infection of *I. mors-panacis* due to the production of high quantities of hydrolytic and oxidative enzymes. The plant is therefore able to successfully retard their invasion. Iron exists on the root surface and within the plant cells and binds to the accumulated phenolic compounds, forming complexes that are likely responsible for the rusty coloration.

Fig. 3. Illustration of ginseng root infection by the highly aggressive *Ilyonectria* species, *I. mors-panacis*, and other weakly aggressive *Ilyonectria* species, leading to the occurrence of rotting and rusty symptoms, respectively. *I. mors-panacis* produces large amounts of cellulases and pectinases, and the plant rapidly responds to invasion by the production of phenolic compounds. However, the pathogen suppresses the toxicity of the phenolic compounds by producing polyphenol oxidases. In parallel, the pathogen sequesters plant and rhizospheric iron by producing siderophores. The other *Ilyonectria* species cannot penetrate the host plant as rapidly as *I. mors-panacis* because they produce lower amounts of hydrolytic and oxidative enzymes. The plant is therefore able to successfully retard their invasion. Iron exists on the root surface and within the plant cells and binds to the accumulated phenolic compounds, forming complexes that are likely responsible for the rusty coloration.

References

[1] Kim YJ, Jeon JN, Jang MG, Oh JY, Kwon WS, Jung SK, Yang DC. Ginsenoside profiles and related gene expression during foliation in Panax ginseng Meyer. J Ginseng Res 2013;38:66–72.
[2] Leung KW, Wong AS. Pharmacology of ginsenosides: a literature review. Chin Med 2010;5:20. http://dx.doi.org/10.1186/1749-6546-5-20.
[3] Wen J, Zimmer EA. Phylogeny and biogeography of *Panax* L. (the ginseng genus, Araliaceae): inferences from ITS sequences of nuclear ribosomal DNA. Mol Phylogenet Evol 1996;6:167–77.
[4] Choi HK, Wen L. A phylogenetic analysis of *Panax* (Araliaceae): Integrating cpDNA restriction site and nuclear rDNA ITS sequence data. Plant Syst Evol 2000;224:109–20.
[5] Attelle AS, Wu JA, Yuan CS. Ginseng pharmacology: multiple constituents and multiple actions. Biochem Pharmacol 1999;58:1685–93.
[6] Kennedy DO, Scholey AB. Ginseng: potential for the enhancement of cognitive performance and mood. Pharmacol Biochem Behav 2003;75:687–700.
[7] Lu JM, Yao Q, Chen C. Ginseng compounds: an update on their molecular mechanisms and medical applications. Curr Vasc Pharmacol 2009;7:293–302.
[8] Proctor JTA. Bailey WG. Ginseng: Industry, botany and culture. Hortic Rev 1987;9:187–236.
[9] Ohh SH, Yu YH, Kim KH, Cho DH. Studies on control of soil-borne diseases and insects of ginseng and development of antifungal compound. In: Ginseng
- Brayford D, Samuels GJ. Some didymosporous species of Chaverri P, Salgado C, Hirooka Y, Rossman AY, Samuels GJ. Delimitation of
- Booth C. Studies of pyrenomycetes. IV.
- Rossman AY, Samuels GJ, Rogerson CT, Lowen R. Genera of the Cabral A, Groenewald JZ, Rego C, Oliveira H, Crous PW.
- Brayford D, Honda BM, Mantiri FR, Samuels GJ. Delimitation of
- Kim YC, Lee JH, Bae YS, Sohn BK, Park SK. Development of effective
- Darmono TW, Owen ML, Parke JL. Isolation and pathogenicity of
- Punja ZK, Wan A, Goswami RS. Root rot and distortion of ginseng seedling
- Chung HS. Ginseng disease. Research reports of the Korean Society of Plant
- Hildebrand AA. Root rot of ginseng in Ontario caused by members of the
- Reeleder RD, Roy R, Capell B. Seed and root rots of ginseng (Panax quinquefolius) in Oregon and Washington, USA. Plant Dis 1993;77:415–8. 
- Shaw K, Kumar RG, Verma S, Dubey RS. Effect of cadmium on lipid peroxi-
- Schroers H-J, Gräfenhan T, Nirenberg HI, Seifert KA. A revision of
- Hemetsberger C, Herrberger C, Zechmann B, Hillmer M, Doehlemann G. The
- Hankins A. Producing and Marketing Wild Simulated Ginseng in Forest and Agroforestry Systems. Produced by Communications and Marketing, College of Agriculture and Life Sciences, Virginia Polytechnic Institute and State University; 2009.
- Rahman M, Punja ZK. Influence of iron on Cyllindrocarpon root rot develop-
- Hankins A. Producing and Marketing Wild Simulated Ginseng in Forest and Agroforestry Systems. Produced by Communications and Marketing, College of Agriculture and Life Sciences, Virginia Polytechnic Institute and State University; 2009.
- Rahman M, Punja ZK. Influence of iron on Cyllindrocarpon root rot develop-
- Chang HS, Bae HW. Ginseng anthracnose in Korea: Factors affecting primary infection, growth of the pathogen, disease development and control. Korean J Plant Prot 1979;18:35–41. 
- Darmono TW, Owen ML, Parke JL. Isolation and pathogenicity of
- Punja ZK, Wan A, Goswami RS, Verma N, Rahman M, Barasubhyte T, Seifert KA, Lévesque CA. Diversity of Fusarium species associated with discolored ginseng roots in British Columbia. Can J Plant Pathol 2007;29:340–53. 
- Reeleder RD, Hoke SMT, Zhang Y. Rusted root of ginseng (Panax quinquefolius) caused by a species of Rhinocerosporidium. Phytopathology 2006;96:1243–54. 
- Choi JE, Ryuik JA, Kim JH, Choi CH, Chon JS, Kim YJ, Lee HB. Identification of endophytic bacteria isolated from rusty-colored root of Korean ginseng (Panax ginseng) and its induction. Korean J Crop Sci 2005;13:1–5. 
- Pathrose B, Jones EE, Jaspers MV, Ridgway HJ. High genotypic and virulence diversity in fuscotorrhiza liroenderi isolates associated with black foot disease in New Zealand vineyards. Plant Pathol 2014;63:613–24. 
- Cymbre WM, Cromble H, Green JB, Lucas J. Pathogenicity of the all fungus to oats: its relationship to the concentration and detoxification of the four avenacins. Phytochemistry 1986;25:2075–83.
- Jewell GC, Roddick JC. Interactive antifungal activity of the glycoalkaloid solanine and chaconine. Phytochemistry 1993;33:323–8. 
- Morisseys JP, Osbourne AE. Fungal resistance to plant antibiotics as a mecha-
- Nicol RW, Traquair JA, Bernards MA. Saponins as host resistance factors in
- Yousef LF, Bernards MA. In vitro metabolism of ginsenosides by the ginseng root pathogen Pythium irregular. Phytochemistry 2006;76(16):1740–9.
- Ivanov DA, Bernards MA. Ginsenosides and the pathogenicity of Pythium irregular. Phytochemistry 2012;78:44–53.
- Shah K, Kumar RG, Verma S, Dubey RS. Effect of cadmium on lipid peroxi-
- Sharma P, Dubey RS. Involvement of oxidative stress and role of antioxidative defense system in growing rice seedlings exposed to toxic concentrations of aluminum. Plant Cell Rep 2007;26:2027–38. 
- Sharma P, Jha AB, Dubey RS. Peroxidase activity. PLoS Pathog 2012;8:e1002684.

The citations and references are not included in the natural text but are part of the document.