Traditional and Molecular Studies of the Plant Pathogen Phytophthora capsici: A Review

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Abstract

Phytophthora capsici is a pathogen that limits the production of diverse crops of economic interest, mainly of the Solanaceae and Cucurbitaceae families. Many studies of this pathogen have been carried out due to its impact on agricultural production systems. This review covers studies on topics such as isolation methods, fungicide resistance, pathogenicity and virulence, physiological races, genetic diversity of its populations, and studies of the genetic expression during the plant-pathogen interaction. The objective of this review is to abridge the information generated on these topics in order to guide future research on this pathogen.

Keywords: Effector; Fungicide; Hemibiotroph; Oomycete; Pathogenicity; Virulence

Introduction

Phytophthora capsici is responsible for large worldwide economic losses and is considered one of the limiting factors in the production of many crops [1]. More than 50 vegetable species have been identified as hosts of this plant pathogen [1,2]. Recently, two leguminous species, Phaseolus vulgaris [3], and Phaseolus lunatus [4], have been identified as hosts of this oomycete. Losses of up to 100% of the crop occur in fields due to the incidence of this pathogen [5].

In Mexico, like in the rest of the world, chili wilt caused by P. capsici is a serious problem. The presence of this soil borne pathogen has been reported in the states of Aguascalientes, Chihuahua, Oaxaca, Puebla, Querétaro, State of Mexico, and Zacatecas [6-12]. The virulence of the isolates of P. capsici found in Mexican fields does not follow a defined distribution pattern and isolates of different levels of virulence can co-exist within the same space [13].

Due to its agronomic importance, numerous studies have been carried out about this plant pathogen which address its isolation [14], morphological characteristics [15], virulence [16], physiological races [17], fungicide resistance [18], the genetic diversity of its populations [19], and its genetic expression during the plant-pathogen interaction [20]. The objective of this review is to abridge the information that has been generated on these topics in order to guide future research on this pathogen.

Isolation and Morphologic Characterization

It is possible to isolate P. capsici from plant tissues or soil samples by using selective media such as PARP (pimaricin, ampicillin, rifampicin, and pentachloronitrobenzene) [21], NARPH (nistatin, ampicillin, rifampicin, pentachloronitrobenzene, and hymexazol) [22], BARP (benomyl, ampicillin, rifampicin, and pentachloronitrobenzene) [23], or with supplemented carrot medium (penicillin, methyl benzimidazol-2-carbamate, pentachloronitrobenzene, and rifampicin) [24].

P. capsici has also been reported in irrigation water. As a model to isolate P. capsici from water both pear and cucumber fruits have been used as baits. The fruits are placed in the irrigation water for intervals of 3 to 7 days. Those portions of the fruits that manifest wounds are transferred to a culture media supplemented with ampicillin and rifampicin [14]. In a similar procedure, pear and eggplant fruits can be used as baits and the wounded parts of the fruits are cultured on water agar media with rifampicin and ampicillin. However, placing the samples of the wounded fruits on the stems of susceptible pepper plants can increase isolation yields. This method can increase isolation percentage between 53%-77% depending on the baiting fruit used [21]. Once the isolation has been carried out successfully, the isolates can be maintained on media such as cotton meal agar, potato dextrose agar, V8 agar, oat agar, rye B agar, or carrot agar [24-28].

Isolating directly from susceptible hosts is the ideal situation as this helps to confirm that the symptoms observed are caused by the pathogen. However, monitoring soil samples and irrigation water can help evaluate the epidemiology of the disease which can thus allow a better management of the pathogen.

Morphological studies of P. capsici isolates recovered from natural populations have shown a high degree of variability respecting the size of the sporangia, pedicels, and oospores [29]. The presence of chlamydospores has been reported on some isolates [5]. Other variable characteristics among isolates are the form of the colony, the form of the sporangia, the abundance of sporangia produced, compatibility type, and optimal growing temperature [15,24,25,30-34]. The colony form can be determined as cottony, roaceous, petaloid, or stellate [25]. The compatibility types (A1 and A2) of each isolate can be known by...
co-culturing each isolate with other isolates of known compatibility type and observing the presence of oospores [35]. Homothallic isolates have been occasionally reported. The presence of both compatibility types on the same field is required so that *P. capsici* can complete its sexual reproductive cycle [1].

Phenotypic variance is one of the first indicators of variability within a population. Of special interest to the study of *P. capsici* is to determine if both compatibility types are present within a population. The presence of both types would indicate the possibility of sexual reproduction and the appearance of new lineages within the population. Sexually reproductive populations implicate different management given that sexual reproduction makes it possible to fix certain traits in the population that may be advantageous to the pathogen, such as fungicide resistance.

**Fungicide resistance**

Oomycetes possess genome plasticity that allows them to develop fungicide resistance [36]. Such is the case of metalaxyl, a systemic fungicide that interferes with the incorporation of uridine during RNA synthesis [37]. The resistance of *P. capsici* to metalaxyl and mefenoxam has been studied in laboratory and field conditions [38,39]. Most studies of *P. capsici* indicate the level of susceptibility of the isolates used to metalaxyl [15-16,21,22,34]. It has been shown that in vitro tests for susceptibility at 100 μg/ml can reliably predict resistance in the field [40].

Fungicide resistance in *P. capsici* has been formally studied for at least two decades (Table 1). When *P. capsici* populations show resistance to mefenoxam and metalaxyl, it is possible to use novel fungicides such as oxamyl, fludioxynil, amitrole, mancozeb, and cymoxanil in alternation to prevent the appearance of resistant isolates [5]. The European and Mediterranean Plant Protection Organization recommends the application of the fungicides etridiazole, propamocarb, chlorothalonil, and copper oxychloride [41]. The effects of copper on the development and infection capacity of different *Phytophthora* species has been extensively studied and shown to be an effective fungicide [42-44].

Mandipropamid and dimetopmorph are mandelamide type fungicides that inhibit the synthesis of lipids and membranes as well as the synthesis of cellulose and cell walls [45-47]. Cyazofamid is a cyanoimidazole type fungicide that inhibits cellular respiration at the complex III of the electron transport chain [48]. Continuous in vitro exposition to dimetomorph has resulted in resistant mutants [49]. Likewise, fludioxonil, flumorph, and pyrimorph resistant mutants have been reported [50-52]. Isolates resistant to pyrimorph present the mutation Q1077K in their cellulose synthase (CesA3) gene [52]. These studies suggest the possibility of generating resistant isolates to these novel fungicides if continuous exposure occurs in the field.

Management of *P. capsici* requires an appropriate use of fungicides. Determining which fungicides are effective against isolates of a given population is important to establish a correct use of such fungicides. However, constant application of one fungicide can lead to the development of resistance within the population. Therefore, special consideration must be taken on which fungicides to apply and at what intervals prevents this from happening.

**Pathogenicity and virulence**

The pathogenicity and virulence of *P. capsici* has been reported for Solanaceous, Cucurbitaceous, and Leguminous species of economic interest such as chili pepper, tomato, cucumber, squash, pumpkin, snap bean, and for Fraser fir and weeds associated with crop fields [2,3,7,15-17,22,24,27,34,53-65]. Inoculation can be done with zoospores [21,64,66] or mycelia [2,16,53]. The capacity of infection is affected by the amount of inoculum, the higher the concentration of the inoculum the higher the aggressiveness of the infection [58,67,68]. When an isolate of *P. capsici* is obtained from a new host it is necessary to confirm its pathogenicity towards the host and corroborate Koch’s postulates. The pathogenicity of *P. capsici* isolates obtained from the roots of *Geranium carthamum*, *Solanum americanum*, and *Portulaca oleracea* were evaluated on chili plants, *S. nigrum*, *S. carolinense*, and *S. capsicoides*. Only *S. nigrum* presented mortality when infected by the pathogen [22]. Isolates of *P. capsici* have also been recovered from snap bean fields (*Phaselus vulgaris*) and their pathogenicity has been confirmed on cultivars of *P. vulgaris* and *P. lunatus*, as well as soy plants (*Glycine max*) [3]. The susceptibility of Fraser fir to *P. capsici* has also been documented [2]. *Arabidopsis thaliana* has been successfully used as an experimental host [69].

Ristaino carried out one of the key studies on *P. capsici* in 1990 [29]. She evaluated the virulence of isolates obtained from seven pepper fields and seven cucurbits fields in North Carolina. She found that some isolates obtained from chili and cucurbits were highly virulent against chili, but that other isolates from cucurbits were less virulent. Thus, this study made it clear that not all isolates have the same virulence level and that the host from which the isolates are obtained may play a role in their virulence towards other hosts.

Temperature, humidity, and the age of the fruit can affect the infection capacity of *P. capsici* over cucumber [55,57] and pepper fruits [70]. The disease is most severe at 25 °C, 4 days after the inoculation. In general, higher temperature and humidity increase the infection capacity of the pathogen. However, ripened fruits are less susceptible. Granke and Hausbeck [71] evaluated the effect of temperature, inoculum concentration, and zoospore age (measured as the number of days after being released from the sporangia) of *P. capsici* on the infection of cucumber fruits. They were susceptible to infection by zoospore suspensions that had a temperature between 9 and 32 °C. The highest infection capacity was exhibited when applying zoospores with concentrations equal to or higher than 5 x 10³ zoospores/ml and temperatures higher than 12 °C. Zoospores of up to a maximum of 5 days of age were able to infect the fruits.

The virulence of a worldwide collection of *P. capsici* isolates was tested on zuccinni, tomato, and chili peppers, by inoculating the fruits with V8 agar discs infested with mycelia. The diameter of the wound, the diameter of the pathogen’s growth on the fruit, and the sporulation density on the fruit were determined. It was concluded that the host of origin from which the isolates were obtained had an effect over the capacity of infection of the isolates on the fruits of other hosts [72].

Virulence and pathogenicity of *P. capsici* has been studied worldwide on a wide range of crops in an effort to determine the interaction between isolates of this pathogen and its wide range of hosts. Virulence and pathogenicity tests are also important to local growers and breeders. These tests allow them to make decisions on which crops are better suited for their local environment and to develop breeding programs to generate resistant or tolerant crops. However, breeding resistant crops to *P. capsici* is difficult due to the presence of physiological races within isolates of this pathogen.

**P. capsici** physiological races

The specificity of plant-pathogen interactions is determined by the host and pathogen genotype. This specificity is easier to study on
plants that have been subjected to genetic improvement, these plants are known as cultivars. Species can present cultivars that are susceptible to a pathogen and cultivars that are resistant to the same pathogen [73]. This phenomenon where isolates of a pathogen have different infection capacities towards diverse cultivars of the same host is known as physiological races and was described by Stakman in 1913 [74].

In order to determine the physiological races of a pathogen it is necessary to determine its interaction with a series of differential hosts [63]. The minimum recommended host age to determine susceptibility to *P. capsici* is when plants have 4 true leaves [66]. *P. capsici* can cause multiple syndromes as it can infect roots, foliage, stems, and fruits [17]. Investigations indicate that different genetic mechanisms are responsible for the resistances to root rot, crown rot, foliar blight, and fruit rot [75]. This allows for the possibility to find physiological races of *P. capsici* for each of the syndromes mentioned in different host species. The studies that have been carried out about *P. capsici* physiological races are listed in Table 2. There are two types of studies: those in which commercial cultivars have been used as differential hosts, and those where the "New Mexico Recombinant Inbreed Lines" (NM-RIL) chili pepper lines have been used. It is also noteworthy that in all of the studies only one Mexican isolate has been used [60].

| Fungicide                          | Test conditions | Concentration range | Inhibition concentration | Reference  |
|------------------------------------|----------------|---------------------|--------------------------|------------|
| 2-(Thiocyanomethylthio)benzothiazole | *in vitro*     | 2-4 g a.i./l         | NA†                      | [8]        |
| Azoxystrobin                       | *in vitro*     | 0.125-4 μg/ml        | 1.92-3.51 μg/ml          | [105]      |
| Copper hydroxide                   | Field          | 1.21 Kg a.i./ha      | NA                       | [18]       |
| Copper sulfate                      | Field          | 30 μg/ml             | NA                       | [15]       |
| Cyclonamid                         | *in vitro*     | 0.1-1000 μg/ml       | >500 μg/ml               | [108]      |
| Cytoxanil                          | *in vitro*     | 0.1-80 μg/ml         | 0.389-8.9 μg/ml          | [105]      |
| Dimethomorph                        | *in vitro*     | 0.1-2.5 μg/ml        | 0.24 μg/ml               | [108]      |
|                                      | Field          | 0.05-1.6 μg/ml       | 0.301-0.365 μg/ml        | [105]      |
|                                      | *in vitro*     | 0.1-0.3 μg/ml        | 0.122-0.203 μg/ml        | [49]       |
|                                      | Growth chamber | 150-600 μg/ml        | 600 μg/ml                | [49]       |
|                                      | *in vitro*     | 0.1-1.0 μg/ml        | 1.02 μg/ml               | [34]       |
| Ethabuxam                           | *in vitro*     | 0.01-100 mg/l        | 1-5 mg/l                 | [26]       |
|                                      | Growth chamber | 32-250 mg/l          | NA                       | [26]       |
| Famoxadone/cymoxanil                | Field          | 0.28 Kg a.i./ha      | NA                       | [18]       |
| Fenamidone                          | Field          | 0.19 Kg a.i./ha      | NA                       | [18]       |
| Flumorph                            | *in vitro*     | 0.05-1.6 μg/ml       | 0.416-0.602 μg/ml        | [105]      |
|                                      | *in vitro*     | 0.5-200 μg/ml        | 0.716-1.363 μg/ml        | [51]       |
|                                      | *in vitro*     | 0.2-0.6 μg/ml        | 0.301-0.487 μg/ml        | [49]       |
| Flumicilide                         | *in vitro*     | 0.08-0.30 μg/ml      | 0.08-0.24 μg/ml          | [50]       |
|                                      | Field          | 0.10 Kg a.i./ha      | NA                       | [18]       |
| Fluorotryptophan                    | *in vitro*     | 5-25 μg/ml           | 10 μg/ml                 | [28]       |
| Mecapropamid                        | *in vitro*     | 0.005-0.5 μg/ml      | 0.03 μg/ml               | [108]      |
|                                      | *in vitro*     | 0.01-1.6 μg/ml       | 0.068-0.080 μg/ml        | [105]      |
|                                      | Field          | 0.15 Kg a.i./ha      | NA                       | [18]       |
| Metalaxyl                           | *in vitro*     | 0.1-800 μg/ml        | 0.243-318 μg/ml          | [105]      |
|                                      | *in vitro*     | 2-4 g a.i./l         | NA                       | [8]        |
|                                      | *in vitro*     | 0.1-1000 μg a.i./l   | 1.41-44.6 μg/ml          | [109]      |
|                                      | Field          | 198 Kg a.i./l        | NA                       | [106]      |
|                                      | Field          | 198 Kg a.i./l        | NA                       | [107]      |
|                                      | *in vitro*     | 1-50 μg/ml           | NA                       | [28]       |
| Metam sodium                        | Field          | 6741 l/ha            | NA                       | [106]      |
| Phosphorous acid                    | *in vitro*     | 12.5-200 μg/ml       | NA                       | [28]       |
| Potassium phosphate                 | Field          | 2.47 y 3.71 Kg a.i./ha | NA                       | [18]       |
| Propamocarb                         | *in vitro*     | 0.55-8.8 μg/ml       | 1.69-3.87 μg/ml          | [105]      |
|                                      | *in vitro*     | 100-1600 μg/ml       | 686-1104 μg/ml           | [105]      |
|                                      | *in vitro*     | 2-4 g a.i./l         | NA                       | [8]        |
| Pyrimorph                           | *in vitro*     | 0-4 μg/ml            | 1.4261 μg/ml             | [52]       |
|                                      | *in vitro*     | 0.5-8 μg/ml          | 3.564-0.02 μg/ml         | [105]      |
|                                      | *in vitro*     | 0.5-6 μg/ml          | 0.557-0.944 μg/ml        | [49]       |
| Telone C-17                         | Field          | 3031 l/ha            | NA                       | [107]      |
| Trifloxostrobin                     | *in vitro*     | 0.125-4 μg/ml        | 3.40-4.08 μg/ml          | [105]      |

*a.i.=active ingredient †=concentration not available

Table 1: *P. capsici* fungicide susceptibility studies.

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only study in which this physiological race phenomena has been tested in
field conditions is the one reported by Hwang and Kim [76], the
rest of the studies have been carried out in greenhouse conditions.
Only one study has evaluated the effect of inoculum concentration
on physiological race typing [53]. It was determined that inoculations
with 10,000 or 100,000 zoospores yielded the same physiological races
of Brazilian P. capsici isolates when using the NM-RIL as differential
hosts. Most studies focus on the median value of the disease index to
rate a specific cultivar as resistant or susceptible to a specific isolate.
In addition mortality rate and area under the disease progress curve (AUDPC)
for each of the interactions are sometimes also reported.

Until now, efforts to produce cultivars universally resistant to all
isolates of P. capsici have not been successful. This is in large part due
to the presence of multiple physiological races of this pathogen across
the world. Therefore, breeders are encouraged to develop cultivars
based on tests on isolates that represent physiological races present
in the growing fields of their interest. Breeding universal resistance
into a host may be something unattainable by breeding programs, but
breeding hosts resistant to a determined number of physiological races
can be a more surmountable challenge.

Genetic diversity

Genetic diversity studies allow us to analyze the population
structure of the pathogen and to confirm the possible genetic
exchange among isolates. The confirmation of this phenomenon in
each geographical area where the pathogen thrives is key in order to
implement a successful management program.

Within the first studies to evaluate the genetic diversity of P. capsici
Restriction Fragment Length Polymorphism (RFLP) assays were
carried out for genomic and mitochondrial DNA [77]. However, it
was not possible to correlate the generated patterns with a geographic
region or host of origin. Numerous studies have reported a wide
range of molecular tools to determine the genetic diversity of P. capsici.
Among these are included molecular markers such as Random
Amplified Polymorphic DNA (RAPD) [65], Amplified Fragment
Length Polymorphism (AFLP) [78], and micro satellites or single
sequence repeats (SSR) [79]. One of the advantages of using AFLP is
the large number of markers (50-70) that can be resolved per reaction
[80]. A characterization of 107 oospores obtained from a cross of two
isolates with distinct AFLP genotyping indicated that AFLP marker
segregation was Mendelian [38].

Another genetic marker that can be used to genotype P. capsici
isolates are single nucleotide polymorphisms (SNP) [79], this can be
achieved with high resolution DNA melting analysis (HR-DMA) [19].
The use of this technique allows the construction of multi locus single
nucleotide polymorphism genotype profiles [81]. These profiles, besides
assessing the genetic diversity of a population of P. capsici, generate
information about the population dynamics and possible genetic
material exchange. P. capsici population studies have demonstrated
that SNP profiles match genotyping with 40 AFLP markers [82]. It is
estimated that the P. capsici genome has a SNP substitution rate of one
different polymorphic site for every 40 bases [83].

An alternative to study the genetic structure of P. capsici
populations is to compare the sequence of distinct polymorphic
nuclear and mitochondrial loci such as Cox1, Cox2, Nad1, Nad5,
β-tubulin, EF-1α, Enolase, HSP90, TigA, Ura3, and ITS [22-23]. Protein
patterns obtained by isoelectric focus have also been used to identify
Phytophthora species [84], as taxonomic criteria [33], or to evaluate
P. capsici diversity [15]. These profiles are performed from total mycelia
protein, from specific enzyme systems like the acid phosphatases, or
polymorphic site for every 40 bases [83].

Population genetics studies of P. capsici can help us understand
the dynamics underlying the epidemiology of this pathogen. Higher
diversity in a population indicates a higher probability of sexual
reproduction within the population. Low diversity could indicate
clonal lineages propagation. Genotyping individuals can help track
their origin and explain their dispersal through a given geographical
area. The sequencing of the P. capsici genome [83] has given way to
the development of SNP markers that allow fine scale studies on the
genetic diversity of this organism. The use of this tool can lead to

| Number of differential hosts | Type of differential hosts                  | Number of isolates evaluated | Origen of isolates | Physiological races determined | Reference |
|-----------------------------|--------------------------------------------|------------------------------|--------------------|-------------------------------|-----------|
| 6                           | Commercial pepper and cucurbits             | 49                           | Georgia            | 10 root rot races              | [65]      |
| 26                          | NM-RIL                                     | 20                           | Brazil             | 8 root rot races               | [53]      |
| 26                          | NM-RIL                                     | 12                           | New Mexico, Turkey, Holland, Argentina | 12 foliar blight races | [59]      |
| 31                          | Commercial pepper lines and cultivars       | 4                            | Michigan, USA      | 4 root rot races               | [54]      |
| 42                          | Commercial tomato cultivars and wild relatives | 4                           | Michigan, USA      | 4 root rot races               | [61]      |
| 26                          | NM-RIL                                     | 17                           | New Mexico, California, and Holland | 13 root rot races | [63]      |
| 11                          | Commercial pepper lines and cultivars       | 34                           | California, New Mexico, North Carolina, and Turkey | 14 root rot races | [56]      |
| 8                           | Commercial pepper lines and cultivars       | 5                            | Guangdong, China   | 2 root rot races               | [24]      |
| 18                          | Commercial pepper lines and cultivars       | 10                           | New Mexico, New Jersey, Italy, Korea, and Turkey | 9 root rot races | 5 foliar blight races | [17]      |
| 21                          | Solanaceous, cucurbits and legumes          | 26                           | Italy              | 13 root rot                   | [34]      |
| 10                          | Solanaceous and cucurbits                   | 30                           | Argentina, California, North Carolina, England, Japan, Mexico, New Mexico, Peru, East Virginia | 14 root rot | [60]      |

Table 2: P. capsici physiological races studies.
better management of the disease by allowing us to understand how the pathogen is being propagated and what forces are driving its population structures.

**Evaluation of the genetic expression during the plant-pathogen interaction**

Most functional genomics studies of *Phytophthora* species have been carried out by genetic transformation, heterologous expression systems, gene silencing, or mutagenesis directed gene disruption [87-90]. However, these type of studies focus on only a handful of genes. It is presumed that most of the processes involved in the infection and colonization of hosts are regulated by multiple genes [91,92].

One of the ways to identify the genes involved in a particular development stage of an organism is to study the transcriptional changes that take place during said stage. The identification of these genes has been facilitated by differential identification methods such as genomic DNA or cDNA libraries hybridizations and subtractive hybridizations [93]. The *P. capsici* genome has been recently sequenced [83]. It is now possible to consult a database that includes all of the open reading frames (ORFs) that have been predicted in the genome of this pathogen (http://genome.jgi-psf.org/Phyc11/Phyc11.download.ftp.html).

Eighty-four effector protein encoding genes of the “crinkler” (CRN) family, related with the infection process, have been identified. The expression of these genes was evaluated during the plant-pathogen interaction of *P. capsici* and tomato at 0, 8, 16, 24, 48, and 72 hours after the inoculation with the help of a microarray composed of the sequences reported for the *P. capsici* genome. The expression levels analysis revealed that some the genes are related to the early stages of the infection while others are related to the late stage. The fusion of some of these effector proteins with green fluorescent proteins (GFP) allowed their localization within the plant cell, where they were found to aggregate around the nucleus and sub nuclear structures [20]. This is consistent with what was previously suggested about effector proteins, which are believed to be the ones in charge of deactivating the defense mechanisms of the plant cell [94,95]. The changes in the genetic expression during the different phases of the infection process have been related to the hemibiotrophic lifestyle of *P. capsici*. The modulation of the genetic expression of this oomycete during its interaction with tomato plants has allowed the identification of four classes of RxLR effectors that are expressed during different physiological process and that are related with changes in the infection process [96]. Three cDNA libraries of *P. capsici* obtained from three different physiological states (mycelia, zoospores, and germinated cysts) have been sequenced. The number of genes expressed during each state corresponded to 13,901 for mycelia, 14,633 for zoospores, and 14,695 for cysts. The large difference in expressed genes during each state is attributed to specific genes required for each of the development states, including 98 genes that codify for effector proteins [97]. Another group of genes related with the infection process that have been recently defined are the “Necrosis Inducing Phytophthora Proteins” (NPP), which participate in the necrosis of the plant tissues [98]. Eighteen genes that encode proteins of the NPP family have been identified active in the mycelia of *P. capsici* [99].

The gene phcnlp1 codifies for a Nep1 like protein of 476 amino acids with a predicted mass of 51.75 kDa. It was isolated from the *P. capsici* isolate Phy12. The expression of this gene was evaluated during the interaction of the isolate with pepper leaves where it was observed that the product of this gene induced visible wounds. The same effect was produced in tabaco (*Nicotiana tabacum*) leaves. These results suggest that phcnlp1l3s directly related to the pathogenesis of *P. capsici* [100].

The gene pcpme6 codifies for a 348 amino acid protein of the pectin methylesterase family with a predicted mass of 38.18 kDa. This gene was isolated from a cDNA library of the *P. capsici* isolate SD33. It was demonstrated that the expression of this gene increases during the infection of pepper leaves, degrading the cell walls and producing necrotic lesions [101]. Similarly, the gene Pcpel1 was obtained from a genomic DNA library of the *P. capsici* SD33 isolate. This gene encodes a 410 amino acid pectate lyase of a predicted mass of 43.8 kDa and has a high level of expression during the interaction of the pathogen with pepper leaves [102]. These studies are consistent with what was reported by Jia et al. [103] which demonstrated a correlation between the virulance of isolates of *P. capsici* and the activity of the enzymes polygalacturonase, pectate lyase, and pectin methylesterase of the isolates cultured *in vitro* in the presence of pepper fruits extracts. The expression of pectin methylesterase genes present in *P. capsici* differs according to the host upon which the infection is taking place [104].

While the use of advanced genomics tools such as microarrays and next generation sequencing have allowed the identification of multiple genes that are suspected to be involved in the infection process of *P. capsici*, much work is needed to validate the functions of these genes. In order to truly understand which of these genes play a significant role in determining the susceptibility of a host it is necessary to determine what are the products of this genes, what their function is, how they internalize into the host, and what their target is within the host. Much work is needed in this area before the information being generated can be successfully used for the development of resistant hosts.

**Conclusion**

The studies about *P. capsici* can be categorized in two main groups: those related to aspects of the integrated management of this pathogen (virulence, pathogenicity, and fungicide resistance), and those that allow us to understand the genetic underpinnings of this organism (genetic diversity and genetic expression).

Studies on pathogenicity and virulence allow breeders and growers to select crops that are better suited to their needs. The information generated by these studies can allow us to discern the isolates into specific physiological races which can be studied in greater detail to improve breeding programs in generating resistant cultivars. Studies focused on the identification of physiological races may have an impact on the management of the disease caused by *P. capsici*, as a large number or physiological races coupled with the presence of both compatibility types in the same field could indicate a zone where the management of the disease is more complicated. The presence of both compatibility types could also facilitate the transmission of fungicide resistance genes, which should be considered when administrating fungicides to a field.

Genetic diversity studies help us to understand the population dynamics of this pathogen and its dispersal patterns. This study could also help us to deduce the forces driving the structure of its populations. Studies that evaluate the genetic expression during the plant-pathogen interaction have helped to identify those genes that are directly related to the infection process. These studies have highlighted the importance of effector proteins, which are attributed the capacity to take control over the plant cell during the infection process, and cell wall degrading enzymes. However, much work is needed to validate the numerous
infection related genes that have been identified in *P. capsici* in order use that knowledge to develop resistant crops.

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