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Cultural, Morphological and Pathogenic Variability in the *Colletotrichum capsici* Isolates, Inciting Anthracnose and Fruit rot of Chilli (*Capsicum annuum* L.) in different Agro-Climatic Zones of Kerala, India

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This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

Aims: To study the cultural, morphological and pathogenic variability existing among the pathogen isolates of *Colletotrichum capsici*, inciting anthracnose and fruit rot of chilli in Kerala. **Study Design:** Completely randomized design (CRD).

Place and Duration of Study: Department of Plant Pathology, College of Agriculture (Kerala Agricultural University), Vellayani, Thiruvananthapuram during 2018-2019.

Methodology: A survey was conducted in the five agro-climatic zones (ACZs) of Kerala *viz.*, Northern zone, Central zone, Special problem zone, High range zone and Southern zone to assess the incidence and severity of anthracnose disease. The various symptoms in the surveyed locations were also noted. Pathogen isolates were grown in PDA medium in the laboratory and colony characters (colour and mycelial growth rate of pathogen colonies) were observed. Also, the morphological characters viz., mycelial width, shape and size of conidia, acervular diameter, number and size of setae, appressorial size were recorded.

Results: Maximum anthracnose incidence of 90 per cent and severity of 52.60 per cent were noticed at Regional Agricultural Research Station (RARS), Pilicode (Northern zone). Characteristic leaf spot, fruit rot and die-back symptoms were observed in the survey locations. Pathogen colonies showed different shades of white, off-white to grey turning brown or black with regular or irregular margins and concentric rings of black acervuli in the potato dextrose agar (PDA) medium. The diameter of growth of *C. capsici* isolates ranged from 7.20 to 8.60 cm at 7 days after incubation. All the *C. capsici* isolates initiated hyaline mycelium having a width of 1.73 - 2.36 μ m and produced numerous sickle shaped conidia with a size range of 19.42 - 20.46 μ m × 2.16 - 3.09 μ m. Black coloured, circular or ellipsoidal acervuli (122.14 - 189.08 μ m dia.) had 20 - 46 setae of length 74.13 to 107.30 μ m. Also, brown or black appressoria of size 8.64 - 12.64 μ m × 5.54 - 7.84 μ m were formed in slide culture. Cc3 from College of Agriculture, Vellanikkara was obtained as the most virulent isolate producing a lesion size of 1.13 cm and PDI of 45.33 in the artificially inoculated chilli fruits (var. Vellayani Athulya) at 5 days after inoculation.

Conclusion: All the nine *C. capsici* isolates exhibited variability in their cultural, morphological and pathogenic characteristics. Variability in phytopathogenic fungi helps in identifying novel and sustainable control methods to tackle the diseases incited by them. Understanding pathogen variability provides new insights for developing effective disease management methods. This will help in promoting sustainable agriculture and preventing severe crop losses.

Keywords: Capsicum annuum; chilli anthracnose; Colletotrichum capsici; variability; pathogenicity; disease incidence; disease severity.

1. INTRODUCTION

Chilli is an important commercial spice crop of Solanaceae family, widely cultivated in the tropical and sub-tropical regions of the world. Fruits are the economic products, known for their pungency and fiery flavour. These form a popular, principal ingredient in the Indian gastronomy. The red ripe fruits are found to be rich in vitamins (A, C and E), riboflavin, thiamine, fibre, proteins and minerals. They are also high in capsaicin and β -carotene levels [1].

India is not only the largest producer but also the major consumer and exporter of chillies in the

world, contributing to about 40 per cent of the total global chilli production and 25 per cent of international exports [2]. But chilli production is affected by a number of factors *viz.*, climate change, lack of high yielding cultivars, poor quality seeds, susceptibility to pests and diseases. About 20-40 per cent yield loss is reported due to weeds, pathogens and pests [3]. Among them, anthracnose disease results in crop losses of 10-54 per cent in India [4].

Colletotrichum capsici is the most common pathogen responsible for chilli anthracnose and fruit rot. The fungus affects almost all the above-ground parts of the chilli plant. Various symptoms

associated with the disease include damping off or seedling blight, leaf spots, die-back of branches, anthracnose and rotting of fruits. The typical symptoms on infected fruits are characterized by sunken, necrotic, circular to angular lesions with concentric rings of black acervuli. On severe infections, pink or orange conidial masses occur on the lesions [5]. In India, anthracnose disease is reported to cause yield losses of 10 - 54 per cent [6]. However, pre- and post-harvest losses of more than 80 per cent have been found to be affecting the marketable yield of chilli plants [7,8].

Pathogen identification is important in preventing the consequences due to development of new pathogen variants, which may result in severe crop losses [9]. Also, variability studies in the pathogen are essential in understanding the prevailing disease situations and the possible control measures. Hence, in the present study, we investigated the cultural, morphological and pathogen variability within the *C. capsici* isolates obtained from the various chilli growing areas in different agro-climatic zones of Kerala.

2. MATERIALS AND METHODS

2.1 Survey and Collection of Infected Fruit Samples

A preliminary survey was conducted in the five agro-climatic zones (ACZs) of Kerala during 2018-2019. The survey locations included Northern Zone (College of Agriculture, Padannakad and Regional Agricultural Research Station (RARS), Pilicode of Kasargod district), of Zone Central (College Agriculture, Vellanikkara, Thrissur and RARS, Pattambi, Palakkad), Special Problem Zone (ORARS, Kayamkulam, Alappuzha and RARS, Kumarakom, Kottayam), Southern Zone (College of Agriculture, Vellayani, Thiruvananthapuram and Farmer's field, Kottarakara, Kollam) and High Range Zone (Farmer's field, Idukki and RARS, Ambalavayal, Wayanad). 100 plants were selected from each of the above survey locations. Disease incidence as well as severity were assessed by counting the diseased and healthy fruits and assessing the extent of the infection in fruits.

2.2 Isolation, Purification and Maintenance of the Pathogen

Anthracnose infected chilli fruit samples were collected from the surveyed regions and brought

to laboratory for the isolation of the pathogen. Small bits of lesions, comprising of diseased and healthy portions were sterilized using 0.1 per cent sodium hypochlorite solution for 1 min and washed three times in sterile distilled water. The bits were blot dried and inoculated on to sterilized plates containing Potato Dextrose Agar (PDA) medium under sterile conditions. Later, the plates were incubated at room temperature $(28 \pm 2 \circ C)$ for 3 days. The emerging hyphal tip was inoculated onto prepared PDA slants. Further. the culture was sub-cultured at regular intervals to maintain the pathogen virulence. The procedure was repeated for all the nine isolates of C. capsici obtained from the survey locations.

2.3 Cultural Characterization of *C. capsici* Isolates

Sterile PDA plates were inoculated with 5 mm mycelial discs taken from actively growing 7-day old cultures of *C. capsici* isolates. The inoculated plates were incubated at room temperature ($28 \pm 2 \text{ °C}$) for 7 days. Three replications were maintained for each isolate. Mycelial growth rate of the different *C. capsici* isolates was measured after incubation for 7 days. The experiment was carried out in three replications of each isolate. Further, cultural observations on the appearance, colour and margin of mycelial growth of the isolates were also recorded.

2.4 Morphological Characterization of *C. capsici* Isolates

Morphological characterization of the different C. capsici isolates was studied by slide culture technique. A glass slide together with cover slips was mounted on two supporting glass rods in a Petri plate containing sterile filter paper. The slide culture unit was then sterilized in an autoclave. Further procedures were carried out in a laminar air flow chamber under sterile conditions. 1 per cent agar medium was prepared and solidified in sterile Petri plates. Square-shaped agar blocks were cut out from these plates and placed at the ends of the sterilized glass slides. The pathogen was smeared on the four corners of each agar block and a cover slip was inserted. Humidity in the slide culture unit was ensured by moistening the filter paper with sterile double distilled water. These units were incubated for 48 h at room temperature (28 ± 2 °C). Further, the coverslips were transferred to a sterile glass slide and stained with lactophenol cotton blue stain. These slides were microscopically observed and the

morphological observations on the mycelial width, shape and size of conidia, diameter of acervuli, number and length of setae, and length and width of appressoria of the *C. capsici* isolates were recorded.

2.5 Preparation of Spore Suspension of the *C. capsici* Isolates

Conidial suspension was prepared using 7-dayold *C. capsici* culture. Initially, the culture plates were flooded with sterilized double distilled water and the surface was slowly scraped with sterile fungal loop. The resulting spore suspension was collected and filtered in two layers of sterilized muslin cloth. Further, the concentration of final conidial suspension was adjusted to 1×10^6 conidia ml⁻¹ using a haemocytometer.

2.6 Pathogenicity of *C. capsici* Isolates

The pathogen isolates tested for were pathogenicity by means of detached leaf and fruit method. Leaves, tender, mature and ripe chilli fruits of Vellayani Athulya (susceptible variety) were used for the experiment. Leaves and tender fruits were collected from 60-day old chilli plants whereas, mature and ripe fruits were picked from 90-day old plants. The detached leaves as well as fruits were surface sterilized by wiping with 70 per cent alcohol. Small incisions were induced, using a sterile needle and inoculated with spore suspensions (1 \times 10⁶ conidia ml⁻¹) of the C. capsici isolates. The inoculated leaves and fruits were incubated at room temperature (28 \pm 2 °C) in sterile plastic boxes lined with moist cotton. The days taken for symptom development and lesion size produced were observed regularly at 3, 5, 7, 10 and 15 days after inoculation. Disease severity was also scored for the different C. capsici isolates, based on the standard score chart (0 = healthy; 1 = 1-5 %; 2 = 5-25 %; 3 = 25-50 % and 4 = 50-100 % of fruit area infected) given by [10]. The pathogen was also re-isolated into PDA medium and compared with the original pathogen culture to confirm the Koch's postulates.

2.7 Analysis OF Data

The design of the experiment was completely randomized design (CRD). The data was statistically analysed using GRAPES software developed by Kerala Agricultural University. Further, the data was subjected to analysis of variance (ANOVA) tests and compared with Duncan's multiple range tests (DMRT) at 5 per cent level (p < 0.05).

3. RESULTS

3.1 Northern Zone Recorded the Maximum Disease Incidence and Severity

A survey was carried out in the five ACZs of Kerala namely Northern zone. High range zone. Central zone. Special problem zone and Southern zone during 2018-2019. The incidence of chilli anthracnose varied from 20 to 90 per cent in the different ACZS. Highest disease incidence (DI) was recorded in the Northern zone, with 90 per cent at RARS, Pilicode, Kasaragod and 85 per cent at College of Agriculture, Padannakkad, Kasaragod in Anugraha variety grown in the area. This was followed by 83 per cent in RARS, Ambalavayal, Wayanad of High range zone (var. Anugraha); 74 per cent in RARS, Pattambi, Palakkad of Central zone (var. Jwalamukhi); 68 per cent in College of Agriculture, Vellayani, Thiruvananthapuram of Southern zone (var. Vellayani Athulya) and 55 per cent in College of Agriculture, Vellanikkara, Thrissur of Central zone (var. Anugraha). Lowest DI of 20 per cent was noticed at farmer's field, Kottarakkara, Kollam of Southern zone (local cultivar). Also, ORARS, Kayamkulam, Alappuzha and RARS. Kumarakom. Kottavam belonging to Special problem zone and Farmer's field, Idukki of High range zone recorded lower DI values of 32 per cent (local variety), 43 per cent (var. Ujwala) and 36 per cent (local variety) respectively.

Disease severity of chilli anthracnose in the various surveyed locations belonging to the five ACZs was observed between 23.63 and 52.60 per cent. Maximum PDI of 52.60 was observed at RARS, Pilicode, followed by 47.90 at RARS, Ambalavayal, 43.20 at College of Agriculture, Padannakkad and 40.33 at RARS, Pattambi. ORARS, Kayamkulam recorded the lowest anthracnose disease severity of 23.63 per cent. Relatively lower PDI values of 36.38, 29.88, 28.25, 27.25 and 24.13 were observed at College of Agriculture, Vellayani; Farmer's field, Idukki; RARS, Kumarakom; Farmer's field, Kottarakkara; and College of Agriculture, Vellanikkara respectively (Table 1).

| Zones | Locations | Variety | *DI | **PDI | Isolates |
|----------|---------------------------------------|---------------|-----|-------|----------|
| | | | (%) | (%) | |
| Northern | College of Agriculture, Padannakad | Anugraha | 85 | 43.20 | Cc1 |
| Zone | (Kasargod) | - | | | |
| | RARS, Pilicode (Kasargod) | Anugraha | 90 | 52.60 | Cc2 |
| Central | College of Agriculture, Vellanikkara | Anugraha | 55 | 24.13 | Cc3 |
| Zone | (Thrissur) | - | | | |
| | RARS, Pattambi (Palakkad) | Jwalamukhi | 74 | 40.33 | Cc4 |
| Special | ORARS, Kayamkulam (Alappuzha) | Local variety | 32 | 23.63 | Cc5 |
| Problem | RARS, Kumarakom (Kottayam) | Ujwala | 43 | 28.25 | Cc6 |
| Zone | | | | | |
| Southern | College of Agriculture, Vellayani | Vellayani | 68 | 36.38 | Cc7 |
| Zone | (Thiruvananthapuram) | Athulya | | | |
| | Farmer's field, Kottarakkara (Kollam) | Local variety | 20 | 27.25 | Cc8 |
| High | Farmer's field, Idukki | Local variety | 36 | 29.88 | Cc9 |
| range | RARS, Ambalavayal (Wayanad) | Anugraha | 83 | 47.90 | Cc10 |
| Zone | | 5 | | | |

Table 1. Incidence and severity of anthracnose and fruit rot of chilli at different survey locations in the five agro-climatic zones (ACZs) of Kerala

* DI – Disease incidence; ** PDI – Percentage Disease Index

3.2 Characteristic Anthracnose Symptoms were Observed on the Infected Chilli Plants of Surveyed Locations

The pathogen infected almost all the aboveground parts of chilli plants. Typical anthracnose symptoms on leaves were noticed as small, water-soaked necrotic lesions, surrounded by yellow halo (Fig. 1A). Fruit rot

symptoms were characterized by water-soaked, circular or elongated, yellowish-brown lesions on the fruit surface. Black acervuli were found concentric scattered or in fashion on the lesions (Fig. 1B). Anthracnose infection on the twigs or branches appeared as brown lesions which advanced from the tip downwards. Severe infection caused defoliation and drying up of affected branches and twigs (Fig. 1C).



Fig. 1. Different symptoms observed on the anthracnose infected chilli plants of surveyed locations (A) The photograph shows minute, water-soaked, dark brown circular or irregular necrotic lesions on the anthracnose infected chilli leaves; (B) Typical fruit rot symptoms depicting greyish brown lesions on the fruit surface, enlarging to cover the entire fruit resulting in drying up and mummification. Severe infection shows concentric rings of acervuli on lesions

3.3 *C. capsici* Isolates Significantly Varied in their Cultural Characteristics on the PDA Medium

Nine C. capsici isolates obtained from the surveyed locations of the five ACZs were plated on PDA plates. Whitish to grey coloured mycelium was initiated; and on microscopic examination, hyaline and septate mycelium produced aseptate, hyaline, unbranched and short conidiophores. The conidia appeared hyaline, one-celled without any septation, sickleshaped with an oil globule in the centre. Nine pathogen isolates (Cc1 - Cc9) were identified as C. capsici while, the isolate from RARS. Ambalavaval (Cc10) was identified as Colletotrichum gloeosporioides. Further, the pure cultures of C. capsici isolates were maintained at lab condition followed by regular sub-culturing at bimonthly intervals.

The *C. capsici* obtained from the different ACZs varied in their cultural characters. The isolates produced sparse mycelium on the PDA medium and concentric rings of black acervuli were observed in the culture. All the *C. capsici* isolates exhibited variation in the colony colour. The colonies varied from white to off-white turning grey, dark brown and black in colour, both on the front as well as rear sides of the culture plates.

The upper side of the culture plates inoculated with the isolates Cc1 of Northern zone, Cc6 of Special problem zone and Cc8 of Southern zone had white mycelium which later turned grey, while the reverse sides exhibited varying degrees of different colours viz., white turning grey, vellowish brown and white turning brown. Cc2 isolate appeared off-white on the front side and vellowish brown in colour on the rear side. In contrast, the isolates, Cc3 and Cc5 produced grey to dark grey mycelium on the PDA medium and their reverse sides exhibited colours of white turning brown and dark brown to black respectively. Cc4 isolate initiated creamy white mycelium on the front side of culture plate and brown to black colour on the reverse side. Cc7 showed greyish white mycelium on the upper side of culture plate while, the lower side exhibited white to greyish white mycelium. However, grey to brown colonies were produced by Cc9 on the upper side compared to dark brown to black on the reverse side (Fig. 2 and Table 2).

The colony margins were observed as either regular or irregular. Seven *C. capsici* isolates *viz.*, Cc2, Cc3, Cc4, Cc5, Cc6, Cc7 and Cc8 displayed regular margins whereas, the isolates Cc1 and Cc9 exhibited irregular margins on the PDA medium.

The *C. capsici* isolates showed significant variation in the growth rate on inoculation in the PDA medium. The isolate Cc4 grew faster in the medium, taking only 7 days to completely cover the Petri plate. This was followed by 8 days for isolates Cc3 and Cc6; 10 days for isolates Cc1 and Cc9 and 11 days for isolates Cc2, Cc5, Cc7 and Cc8. About 7 days after incubation, highest growth of 8.6 cm was observed for the isolate Cc4 followed by 8.5 cm in Cc6 and 8.2 cm in Cc3. In contrast, minimum growth of 7.2 cm was observed for the isolate Cc2 (Table 2). These plates produced orange to pink coloured spores in culture, on continued storage for about 20 to 25 days.

3.4 Morphological Variability Depicted Remarkable Variation in the *C. capsici* Isolates

Microscopic observations of mycelia, conidia, setae, acervuli and appressoria of nine C. capsici isolates were recorded on the PDA medium. Width of mycelia in C. capsici isolates varied between 1.73 µm and 2.39 µm. Highest mycelial width was observed in isolate Cc5 of Special problem zone (2.39 µm), while the minimum was recorded in Cc2 of Northern zone (1.73 µm). All isolates produced single-celled, sickle the shaped conidia having a central oil globule. Conidial length varied between 19.42 µm and 20.46 µm whereas, the width of conidia ranged from 2.16 µm to 3.09 µm. Maximum length of conidia was observed in Cc4 isolate of Central zone (22.68 µm) while, conidial length was minimum for Cc5 of Special problem zone (19.42 um). In contrast, Cc8 isolate from Southern zone produced conidia having maximum width of 3.09 µm and lowest conidial width was seen in the isolate Cc1 of Northern zone (2.16 µm) (Table 3).

The *C. capsici* isolates gave rise to black, round or elliptical acervuli having diameter ranging from 122.14 μ m to 189.08 μ m. The isolate Cc3 of Central zone produced acervuli with highest diameter of 189.08 μ m whereas minimum acervular diameter of 122.14 μ m was recorded in Cc8 isolated from Southern zone.

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Fig. 2. Colony characters of the nine *C. capsici* isolates on PDA medium at 7 days after incubation. depicting the upper as well as reverse side of the culture plates (A) Photographs show the upper side of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represents the rear view of the culture plates inoculated with the nine *C. capsici* isolates numbered from Cc1 to Cc9; (B) represe

| Table 2. Cultural characteristics of the unreference. <i>Capsici</i> isolates on FDA medium |
|---|
|---|

| Isolates | Appearance | Colour of mycelia | | Margin | DTCP | Radial growth of | |
|----------|---------------------|--------------------|---------------------|-----------|------|-----------------------------|--|
| | | Upper side | Reverse side | | | pathogen mycelium (cm) | |
| Cc1 | Sparse mycelial | White turning grey | White turning brown | Irregular | 10 | $7.8 \pm 0.24^{\circ}$ | |
| Cc2 | concentric rings of | Off-white | Yellowish brown | Regular | 11 | 7.2 ± 0.22^{e} | |
| Cc3 | acervuli | Grey to dark grey | White turning brown | Regular | 8 | 8.2 ± 0.26^{b} | |
| Cc4 | | Creamy white | Brown to black | Irregular | 7 | 8.6 ± 0.17^{a} | |
| Cc5 | | Grey to dark grey | Dark brown to black | Regular | 11 | 7.3 ± 0.08^{de} | |
| Cc6 | | White turning grey | Yellowish brown | Regular | 8 | ^{ab} 8.5 ± 0.13 | |
| Cc7 | | Greyish white | White turning brown | Regular | 11 | 7.4 ± 0.15 ^{de} | |
| Cc8 | | White turning grey | White turning black | Regular | 11 | 7.4 ± 0.22^{d} | |
| Cc9 | | Grey to brown | Dark brown to black | Irregular | 10 | 7.6 ± 0.21^{cd} | |
| SE(m) ± | | | | | | 0.037 | |
| CD(0.05) | | | | | | 0.281 | |

DTCP - Days taken to completely cover Petri dish

| Isolates | Width of | | Conidia | | Diameter of |
|----------|---------------------------|----------------|---------------------------|--------------------------|----------------------------|
| | Mycelia (µm) | Shape | Length (µm) | Width (µm) | acervuli (µm) |
| Cc1 | 1.83 ± 0.37 ^{cd} | Sickle shaped | 21.46 ± 1.24 ^b | 2.16 ± 0.42 ^e | 146.26 ± 3.14 ^e |
| Cc2 | 1.78 ± 0.39^{d} | conidia with a | 21.20 ± 1.02 ^b | 2.30 ± 0.49^{de} | 142.19 ± 2.08^{f} |
| Cc3 | 1.86 ± 0.11^{cd} | central oil | 20.46 ± 3.57^{bcd} | 3.00 ± 0.38^{a} | 189.08 ± 2.58^{a} |
| Cc4 | 2.03 ± 0.13^{bcd} | giobule | 22.68 ± 1.38 [°] | 2.32 ± 0.51^{cde} | $173.02 \pm 2.57^{\circ}$ |
| Cc5 | 2.39 ± 0.24^{a} | | 19.42 ± 1.87 ^d | 2.52 ± 0.34^{bc} | 143.02 ± 1.63^{1} |
| Cc6 | 2.05 ± 0.24^{bcd} | | 20.52 ± 0.85^{bc} | 2.40 ± 0.20^{bcd} | 153.31 ± 4.09^{d} |
| Cc7 | 2.12 ± 0.11^{abc} | | 21.83 ± 0.59^{bc} | 3.02 ± 0.42^{a} | 128.89 ± 1.70^{9} |
| Cc8 | 2.21 ± 0.23 ^{ab} | | 19.90 ± 3.68^{cd} | 3.09 ± 0.46^{a} | 122.14 ± 1.47^{h} |
| Cc9 | 2.25 ± 0.15^{ab} | | 20.57 ± 2.09^{bc} | $2.57 \pm 0.43^{\circ}$ | $167.97 \pm 1.42^{\circ}$ |
| SE(m) ± | 0.06 | | 4.41 | 0.17 | 5.98 |
| CD | 0.31 | | 1.06 | 0.21 | 2.18 |
| (0.05) | | | | | |

Table 3. Morphological characteristics of different C. capsica isolates

A number of long, dark brown to black, needlelike, elongated and septate setae were observed with the acervuli of *C. capsici* isolates, varying from 15 to 26. The number of setae reached a maximum of 30-46 in the acervuli of isolate Cc3 of central zone compared to lowest number of setae in Cc1 isolate of Northern zone (15-36). Setae length in the nine *C. capsici* isolates varied between 74.13 μ m and 107.30 μ m. Maximum length of 107.30 μ m was recorded in the setae produced by the isolate Cc6 (Special problem zone) and minimum of 74.13 μ m in Cc9 of High range zone.

Penetration and attachment of the *C. capsici* isolates was made possible by the production of dark brown, round or elliptical appressoria. The appressorial size varied from 8.64 - 12.64 μ m x 5.54 - 7.84 μ m. Appressorial length and width was highest in the isolate Cc3 of Central zone (12.64 μ m; 7.84 μ m). Minimum length (8.64 μ m) and width (5.54 μ m) was measured for the appressoria initiated by the isolates Cc6 and Cc5 respectively, belonging to the Special problem zone (Table 4).

3.5 Cc3 Isolate was the Most Virulent among the *C. capsici* Isolates

Pathogenicity of *C. capsici* isolates were tested by pin pricking the chilli fruits, followed by inoculation. All the *C. capsici* isolates were pathogenic to chilli and produced anthracnose symptoms on the artificially inoculated fruits of chilli (var. Vellayani Athulya). The days taken for initiating anthracnose symptoms was reduced to 1 day in the *C. capsici* isolates, Cc1 and Cc2 from Northern zone, Cc3 from Central zone and Cc6 from Special problem zone. While, the other *C. capsici* isolates produced the symptoms in 2 to 3 days after inoculation. However, a maximum of 3 days for symptom appearance was recorded in the isolates Cc4 from Central zone, Cc7 and Cc8 from Southern zone.

The isolate Cc3 produced a highest lesion size of 1.13 cm, followed by Cc6 (0.91 cm) and Cc2 (0.73 cm). In contrast, minimum lesion size was recorded in chilli fruits inoculated with Cc8 (0.17 cm). The other isolates Cc1, Cc4, Cc5, Cc7 and Cc9 also formed lesions of 0.50, 0.33, 0.23, 0.28 and 0.37 cm respectively on the 5th day after inoculation.

Maximum disease severity of 45.33 per cent was recorded at 5 days after artificial inoculation of chilli fruits with Cc3, followed by Cc6 (41.67 %) and Cc2 (33.33 %). Lowest disease severity of 16.67 per cent was observed in the fruits, on inoculation with Cc7 and 20.33 per cent due to Cc8. PDI of 25 was recorded in all the other *C. capsici* isolates (Cc1, Cc4, Cc5 and Cc9) (Table 5).

4. DISCUSSION

The survey performed in the five ACZs of Kerala during 2018-2019 revealed the prevalence of *C. capsici* as the major causal agent of chilli anthracnose disease. Further, the disease incidence as well as severity varied significantly

in the surveyed areas of various districts. The difference in incidence and severity of chilli anthracnose existing in the surveyed locations may be accredited to the cultivated variety, unpredictable climatic conditions, virulence of the strains and management strategies involved. Anthracnose incidence in the four districts of Jaipur ranged between 51.75 and 66.70 per cent [11]. Highest anthracnose incidence and PDI of 80 per cent and 54 respectively was observed at RARS, Ambalavayal [12]. An extensive survey in the 36 locations of Uttar Pradesh, where maximum anthracnose severity was recorded at Jaunpur (54.91 %) [13]. [14] conducted a roving survey in the chilli cultivating districts of Karnataka during 2020-2021. Maximum PDI of 36.95 was observed in the Dharwad district compared to lowest in the Belagavi district (32.93).

Characteristic fruit rot symptoms with concentric rings of acervuli and dieback symptoms on infected twigs were also recorded in the survey locations. [15] reported the presence of dark, greyish brown lesions on the anthracnoseinfected stems and leaves, having scattered or concentric rings of acervuli. [16] found the appearance of sunken, circular or round spots, having dark brown to black margins on the infected ripe chilli fruits. [17] observed circular, black coloured, necrotic lesions on unripe fruits. Under severe conditions, black acervuli were found on the lesions, causing fruit decay. Infected branches displayed die-back symptoms.

| Table 4. Setae and appressori | al characteristics of diffe | rent C. capsici isolates |
|-------------------------------|-----------------------------|--------------------------|
| | | |

| Isolates | ç | Setae Appr | | ressoria | |
|-----------|--------|----------------------------|--------------------------------|-------------------------------|--|
| | Number | Length (µm)* | Length (µm) | Width (µm) | |
| Cc1 | 15-36 | 80.92 ± 2.33^{d} | 11.13 ± 0.46 [°] | 5.63 ± 0.62^{ef} | |
| Cc2 | 20-32 | 78.72 ± 3.31^{d} | 11.48 ± 0.69 ^{bc} | ^{def} 5.71 ± 0.81 | |
| Cc3 | 30-46 | 106.37 ± 5.08^{a} | 12.64 ± 0.87^{a} | 7.84 ± 0.57^{a} | |
| Cc4 | 32-45 | 101.98 ± 2.78 ^b | 9.05 ± 0.90^{e} | 6.48 ± 0.41^{bc} | |
| Cc5 | 25-37 | $94.22 \pm 3.30^{\circ}$ | ^{abc} 11.95 ± 0.58 | 5.54 ± 0.64^{f} | |
| Cc6 | 38-43 | 107.30 ± 2.91^{a} | 8.64 ± 1.62 ^e | 6.02 ± 0.21^{cde} | |
| Cc7 | 24-40 | $96.62 \pm 1.85^{\circ}$ | $11.21 \pm 0.32^{\circ}$ | 6.10 ± 0.67^{cd} | |
| Cc8 | 20-30 | $94.60 \pm 1.18^{\circ}$ | 12.33 ± 0.91 | 6.13 ± 0.26^{cd} | |
| Cc9 | 27-40 | $74.13 \pm 0.86^{\circ}$ | 9.96 ± 1.63^{d} | 6.78 ± 0.19^{b} | |
| SE(m) ± | | 8.31 | 0.98 | 0.28 | |
| CD (0.05) | | 2.57 | 0.88 | 0.47 | |

| Table 5. Pathogenicity | y of <i>C. c</i> | <i>apsici</i> isolates o | n mature chilli fru | its of variet | y Vella | yani Athul | ya |
|------------------------|------------------|--------------------------|---------------------|---------------|---------|------------|----|
| | | | | | - | | |

| Isolates | Days taken for symptom appearance | Lesion size on fruits (cm) at 5 th day after inoculation | PDI (%) |
|-----------|--------------------------------------|--|---------|
| Cc1 | 1 | 0.50 ± 0.20^{cd} | 25 |
| Cc2 | 1 | 0.73 ± 0.05^{bc} | 33.33 |
| Cc3 | 1 | 1.13 ± 0.20ª | 45.33 |
| Cc4 | 3 | 0.33 ± 0.15^{de} | 25 |
| Cc5 | 2 | 0.23 ± 0.06^{e} | 25 |
| Cc6 | 1 | 0.91 ± 0.20^{b} | 41.67 |
| Cc7 | 3 | 0.28 ± 0.17^{de} | 16.67 |
| Cc8 | 3 | 0.17 ± 0.06^{e} | 20.33 |
| Cc9 | 2 | 0.37 ± 0.15^{de} | 25 |
| SE(m) ± | | 0.023 | |
| CD (0.05) | | 0.260 | |

Significant variations in terms of cultural and morphological characters were observed among the nine *C. capsici* isolates. The fungal colonies produced on the PDA medium showed different shades of white, off-white to grey turning brown or black. About 7 to 11 days were taken by the isolates to completely cover the PDA plates. The diameter of growth produced by the nine C. capsici isolates also varied between 7.20 and 8.60 cm at 7 days after incubation. Morphological showed observations also remarkable differences among the C. capsici isolates causing chilli anthracnose. Similar results have been reported by several researchers.

Whitish grey mycelium of C. capsici isolates, produced conidia with size 19.70 - 33.60 µm × 2.23 - 4.86 µm [18]. White to grey, sparse to cottony mycelium with regular or irregular margins were found in the C. capsici cultures. Conidial length ranged from 13.50 um to 21.20 um while width varied between 3.20 µm to 4.80 um. Black, circular acervuli measured 130 -162.60 µm in diameter. The length of setae varied between 145.40 µm and 179.10 µm [19]. Similarly, [20] reported that the ten isolates of C. capsici (Tamil Nadu) initiated fluffy to cottony white aerial mycelium having irregular to regular margins in the PDA medium and highest mycelial growth diameter of 8.15 cm was noticed at 7 days after inoculation. All the isolates produced falcate conidia having a centrally placed oil globule and acervuli varying in size (18 - 23 µm × 3.43 - 3.97 µm) and mature acervuli comprised of several setae (12 - 32). [21] recorded fluffy or cottony to felty mycelium produced by C. capsici isolates of Andhra Pradesh with a growth rate of 3.8 to 9.8 mm per day on PDA. Conidial size appeared to be measured between 18.10 - 22.36 µm in length and 2.84 - 4.05 µm in width. The pathogen cultures were observed as grevish to white in colour, having fluffy texture and regular margins. Hyaline, sickle shaped conidia of size ranging between 18 - 27 µm in length and 2.10 -4.10 µm in width was produced in culture. The setae were dark coloured, septate measuring 110 - 272 µm × 4 - 6 µm in size [22]. [23] noticed maximum radial growth in all the nine isolates of C. capsici of Eastern Uttar Pradesh, grown in PDA medium at 9 days after inoculation. These isolates produced cottony or fluffy mycelium with either regular or irregular margins. Conidia were falcate shaped, with a size ranging between

18.10 - 27.10 μ m × 1.6 - 2.3 μ m. Mature acervuli contained many setae with varying length of 87.20 - 151.40 μ m and width of 3.30 - 5.30 μ m.

The differences in morphological as well as the cultural characteristics within the pathogen species may be due to the prevailing temperature, nutritional source and pH of medium. [24] showed that the significant changes observed in colony colour as well as growth rate of C. capsici can be accorded to the nutritional factors present in the media. Significant increase in the conidial size as well as spore volume of the postharvest pathogens, Botrytis allii and Pencillium hirsutum were reported by [25]. This was attributed to the changes in temperatures ranging from 20 to -2°C. In contrast, spore size was noticed minimum at higher temperature and an enhancement in conidial size and spore volume was reported with decreasing temperature of -2°C. Similarly, lower incubation temperatures marked an enhancement in the conidial size of entomopathogenic fungi. Metarrhizium brunneum [26]. Whereas, [27] reported the presence of large sized conidia in the chickpea wilt pathogen, Fusarium oxysporum f. sp. ciceris on exposure to higher temperatures of 30°C, with additions of carbon and nitrogen, at pH of 6 - 7 in the culture medium. Also, recombinations and mutations form remarkable factors in the development of fungal variability. [28] reported the involvement of random matings as well as mutations as a source of genetic variability in phytopathogenic fungi.

The isolates of C. capsici were inoculated on the detached chilli fruits (var. Vellayani Athulya) by spraying spore suspension. Maximum lesion size of 1.13 cm and thus, the maximum disease severity was recorded in the chilli fruits inoculated with the isolate Cc3 of Central zone. Minimum lesion size of 0.17 cm was observed for the isolate Cc8 obtained from Southern zone (Farmer's field, Kottarakkara). However, Cc7 isolate recorded lowest disease severity of 16.67 per cent. [15] categorized the ten isolates of C. capsici (Thailand) into three groups viz., mildly virulent, moderately virulent and severely virulent, based on the disease scores incited in the inoculated fruits. [29] reported that among the twenty C. capsici isolates of Tamil Nadu, Cc1 incited maximum intensities of chilli fruit rot (69.90 %) and leaf rot (63.20 %) infections in artificially inoculated chilli plants. [30] recorded that the C. capsici isolate. UDR Cc-01 to be the most virulent among the four isolates, causing a maximum PDI of 61.20 on the chilli variety, Pusa Jwala. [31] observed the pathogenic variability of ten C. capsici isolates obtained from Rajasthan. The highly virulent isolate was recorded as UDP

Cc1 with a severity of 47.17 per cent while, the least virulent was Raj Cc1 with a PDI of 26.42.

5. CONCLUSION

Survey conducted in the five ACZs of Kerala revealed that RARS, Pilicode of Northern zone (Kasargod district) is more prone to anthracnose infection with 52.60 PDI. Nine C. capsici isolates were obtained from the survey locations and their cultural, morphological as well as pathogenic differences were studied. Among the C. capsici isolates, Cc3 of College of Agriculture, Vellanikkara (Central zone) was obtained as the most virulent isolate. These variability studies comprehend towards the selection of better management options and also reduce the risk of creation of new pathogen races. The integration of different management measures can help in mitigating severe crop losses.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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