

Pathogenicity and Molecular characterization of *Colletotrichum gloeosporioides* causing anthracnose disease in anthurium

ABSTRACT

Anthurium is an excellent cut flower crop commercially cultivated throughout the world. Anthracnose or spadix rot disease leads to massive flower loss to anthurium growing farmers. Survey was conducted for anthurium anthracnose disease during the year 2017-2018 in anthurium growing areas of Tamil Nadu, India. *Colletotrichum gloeosporioides* cultures (10 isolates) were isolated and pathogenicity test was proved by several artificial inoculation methods. Among this, pin prick plus spraying spore suspension method has recorded the highest per cent disease index of 64.38 on anthurium plants. Molecular analysis based on obtained sequences (MH479426) of the rDNA internal transcribed spacers (ITS1 and ITS4) resulted more than 92 per cent identical to ITS sequence of *C. gloeosporioides* published in GenBank database. The molecular study confirmed that anthracnose disease of anthurium is caused by *C. gloeosporioides*. The molecular-based clustering demonstrated the genetic relationships of the isolates and species of *Colletotrichum* and indicated that ITS rDNA sequence data were potentially useful in taxonomic species determination.

Keywords: Virulence, *Colletotrichum gloeosporioides*, ITS, Incidence, phylogenetic, Inoculation

1. INTRODUCTION

Anthurium (*Anthurium andreanum*) is one of the beautiful cut flowers cultivated mostly in tropical humid climate. It is an excellent cut flower crop commercially cultivated throughout the world. The flowers of anthurium are popular among flower arrangers because of their attractive colours, increased vase life, bold effect and long lasting qualities (Bhatt and Desai, 1989). The popularity of growing anthurium as cut flowers has risen tremendously in the past few years and it has now become an important export-oriented crop.

Anthracnose or spadix rot disease caused by *Colletotrichum gloeosporioides* is a major constraint to anthurium production which leads to massive losses in terms of quality and quantity. *Colletotrichum* genus was recently voted as the eighth most important group of plant pathogenic fungi in the world, based on perceived scientific and economic importance (Dean *et al.*, 2012). Anthracnose caused severe rotting incidence of anthurium resulting in 100 per cent death of plants in Alleppy district of Kerala (Santhakumari *et al.*, 2001). Severity of anthracnose in anthurium ranged from 21.67 to 54.89 per cent in Tamil Nadu (Nandinidevi, 2008).

Colletotrichum is one of the most common plant pathogenic fungi in tropical or temperate regions. The symptoms can appear as small, water-soaked speckles or large necrotic and chlorotic lesions that are circular to irregular shape. Lesions are yellow, brown or black in colour, often with a bright-yellow chlorotic halo. This research paper deals about the molecular characterization of *Colletotrichum* species associated with anthracnose disease of anthurium and standardization of inoculation techniques for proving the pathogenicity.

2. MATERIALS AND METHODS

2.1. Survey and disease incidence

49 Survey was conducted in different anthurium growing areas of Tamil Nadu, India during
50 2017–2018 and the disease incidences were recorded. The survey includes the observation of
51 symptoms over time and the Per cent Disease Index was calculated by McKinney (1923)
52 formula. Anthracnose disease infected leaf samples were collected from different anthurium
53 growing areas of Tamil Nadu for isolation of the fungus, *Colletotrichum gloeosporioides*.

54 **2.2. Isolation of *C. gloeosporioides***

55 The diseased leaf was first washed with tap water to remove dust and other contaminants.
56 The infected portion was cut into small bits and surface sterilized with 10 per cent sodium
57 hypochlorite for 5-10 minutes. In order to remove the residue of the chemical, the tissue bits
58 were washed with three changes of sterile distilled water. The surface sterilized bits were
59 placed on Potato Dextrose Agar (PDA) medium in sterilized Petri dishes. These plates were
60 incubated at room temperature ($28 \pm 2^\circ\text{C}$) for seven days. After incubation, the cultures were
61 purified by hyphal tip method (Dhingra and Sinclair, 1985) and the fungal cultures were
62 maintained separately in agar slants/Pert plates.

63 **2.3. Pathogenicity assay**

64 The Pathogenicity test was conducted under protected cultivation. Anthurium plants
65 of healthy temptation variety were maintained in pots under 75% shade. Among the ten
66 isolates, the virulent isolate ICg1 was taken for different inoculation methods. The virulent
67 fungal culture filtrate was prepared and the spore suspension was adjusted to 5×10^6
68 conidia/ml. The different inoculation methods were:

69 **2.3.1. Leaf-clipping method**

70 Three leaves of anthurium plants were clipped off approximately 2-3 cm from their
71 tip by using a pair of scissors dipped in the inoculum of each isolate prepared as described
72 earlier. Similarly control was inoculated with sterile distilled water. The plants were wrapped
73 in moist plastic bags to conserve moisture and placed in greenhouse at $25-27^\circ\text{C}$ immediately
74 after inoculation until optimum disease development (Schaad, 1980).

75 **2.3.2. Pin-pricking method**

76 Needles were dipped into the fungal cell suspension of each isolate prepared and
77 gently prick the leaf blades of anthurium plants maintained in pots. Similarly control was
78 inoculated with sterile distilled water. The plants were covered with moist plastic bags to
79 conserve moisture and placed in greenhouse chamber immediately after inoculation (Klement
80 and Goodman, 1967).

81 **2.3.3. Spray inoculation method**

82 Suspensions of different fungal isolates were sprayed directly onto anthurium plants
83 potted separately for pathogenicity test. Then the treated pots and sterile water treated control
84 pots were incubated for observation. (Mew, 1989).

85 **2.3.4. Injury by carborandom powder**

86 The anthurium plants were artificially injured by carborandom powder randomly.
87 Then the prepared fungal spore suspension of each isolate was sprayed onto the anthurium
88 plant accordingly under protected cultivation. The plants sprayed with sterile water served as
89 control and the plants were observed for disease development.

90 **2.4. DNA extraction and PCR amplification**

91 The different isolates of *Colletotrichum gloeosporioides* were grown in potato dextrose
92 broth separately for fifteen days. The fresh mycelium from the broth was harvested by filtration
93 separately with sterile filter paper and by using liquid nitrogen; it was grinded to a fine powder.
94 100-200 mg of mycelial powder of each isolate was taken separately and macerated with CTAB
95 buffer. After maceration 700 μl solution of each isolate was transferred into centrifuge tube
96 separately and incubated at 65°C for 25 minutes. Equal volume of phenol: chloroform: isoamyl
97 alcohol (25:24:1) were added to the incubated tubes and centrifuged at 10000 rpm for ten

98 minutes. After centrifugation, the aqueous layer from the tubes was separated and chloroform:
99 isoamyl alcohol (24:1) of equal volume was added to precipitate the contaminants.

100 The mixture was again centrifuged for ten minutes at 10000 rpm. After centrifugation 300
101 µl aqueous solution of each isolate was taken separately to which 5M sodium acetate: ice cold
102 ethanol (5:2) was added. This mixture was kept at -20 °C overnight. This mixture was again
103 centrifuged at 13000 rpm at 4 °C for ten minutes. After centrifugation equal volume of ethanol
104 was added, centrifuged and kept for pellet drying. Then 50 µl of TE-buffer was added into the
105 pellet separately, then the genomic DNA of each isolate was checked by Polymerase Chain
106 Reaction (PCR) in order to amplify the DNA by using universal primers ITS1 forward (5'-
107 TCCGTAGGTGAACCTGCGG-3') and ITS4 reverse (5'-TCCTCCGCTTATTGATATGC-3')

108 PCR amplification was done in a thermal cycler. The total volume for PCR reaction was
109 ten µl which contained seven µl PCR master mix (Taq DNA polymerase, AMPLICON
110 Company), one µl of free nuclease water, 0.5 µl of forward (ITS1) and reverse primer (ITS4), and
111 one µl of DNA template. Thermo cycling procedure was carried out at initial denaturation (95°C)
112 for two minutes, followed by 38 cycles of denaturation (95°C) for one minute, annealing at 55°C
113 for 30 seconds, one minute extension at 72 °C and final extension for ten minutes at 72° C.

114 2.5. Sequence analysis

115 DNA sequencing of PCR product was sequenced in both directions (Eurofins
116 Genomics India Pvt Ltd, Bangalore). The expected size of the PCR product was
117 approximately 550bp. The obtained sequences were analysed using BLAST program with
118 default parameters (NCBI) (Altschul *et al.*, 1990). ITS sequences were used to construct two
119 unrooted phylogenetic trees, using the MEGA (version 10.0.2.) program (Kumar *et al.*, 2018)
120 with the neighbour-joining (NJ) method Saitou and Nei, (1987).

121 The statistical reliability of the branches was evaluated by performing bootstrap
122 analysis (1000 replicas). Sequence data matrix was aligned using the ClustalW option and the
123 alignment was corrected where necessary. ITS sequences of the species *C. gloeosporioides*
124 (GenBank accession numbers MH479426) were compared with eleven ITS sequences
125 belonging to *Colletotrichum* sp. showed more than ninety per cent identity.

126 3. RESULTS AND DISCUSSION

127 3.1. Survey and Isolation

128 Survey was conducted during 2017-2018 in various places of Tamil Nadu. The
129 occurrence of anthracnose disease on anthurium cultivars was recorded (Table 1). Symptoms
130 typical of anthracnose caused by *Colletotrichum gloeosporioides* were frequently observed
131 on anthurium plants in various districts of Tamil Nadu, India. In general, the common
132 symptoms consist of necrotic spots on leaves were more or less circular, light brown with a
133 prominent yellow margin that in a later phase of infection formed the fruiting bodies of the
134 fungus. Anthracnose appeared as small, irregular and sunken lesions, with a diameter of 0.5
135 cm or little more, which could turn to black, increase in size and form acervuli.

136 The present survey revealed that the disease incidence of anthracnose on leaves
137 caused by *C. gloeosporioides* was the most serious problem in the cultivation of anthurium in
138 Tamil Nadu, India. Severe symptoms were frequently observed on leaves as well as flowers.
139 This paper explained about the pathogenicity and identification of different pathogenic
140 isolates of *Colletotrichum gloeosporioides* based on molecular characterization.

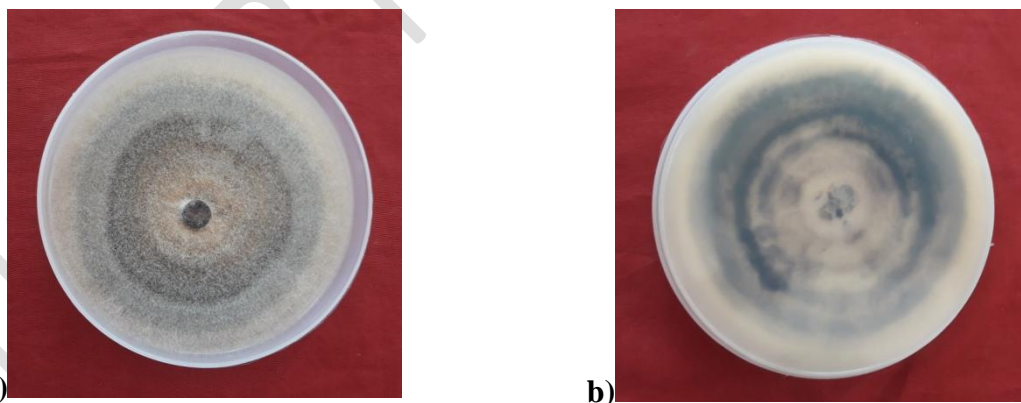
141 A total of ten *C. gloeosporioides* isolates were isolated of which all the ten were
142 identified as *C. gloeosporioides* through morphological and molecular analysis. Colonies
143 grown on PDA were initially white-orange and then turned to greenish-grey (Fig. 1a). In the
144 reverse side, colonies were brownish-orange to grey (Fig. 1b). Bright orange to pink spore
145 masses were produced. Conidia were hyaline, unicellular and cylindrical. The length and
146 width of conidia produced by *C. gloeosporioides* sized from 10 to 16 µm in length and 3 to 5
147 µm in width.

148 **Table 1. Survey and isolation of anthurium anthracnose disease on different anthurium**
 149 **growing areas of Tamil Nadu, India.**

S.No.	Village name	District	Isolate code	Per cent Disease Index (PDI)*	Geographical location		
					Latitude	Longitude	Altitude (Ft.)
1.	Thandikudi	Dindugal	ICg1	56.66 ^a	10° 17' 36.8658"N	77° 42' 33.3324"E	3832.021
2.	Pechipparai	Kanyakumari	ICg2	48.88 ^d	8° 26' 26.145"N	77° 18' 15.7536"E	393.701
3.	Thuckkalay	Kanyakumari	ICg3	38.88 ^f	8° 14' 33.0246"N	77° 18' 53.9382"E	32.8084
4.	Parasalai	Kanyakumari	ICg4	52.21 ^c	8° 19' 37.8834"N	77° 9' 29.7354"E	104.987
5.	Moolachanvilai	Kanyakumari	ICg5	31.10 ^h	8° 12' 14.0106"N	77° 9' 29.7354"E	131.234
6.	Yercaud	Salem	ICg6	31.10 ⁱ	11° 47' 44.8398"N	78° 12' 42.8148"E	4921.26
7.	Nagloor	Salem	ICg7	17.77 ^j	11° 50' 7.0218"N	78° 12' 22.809"E	5324.803
8.	TNAU	Coimbatore	ICg8	36.66 ^g	11° 0' 56.0298"N	76° 55' 51.312"E	1430.45
9.	Pandrimalai	Dindugal	ICg9	54.43 ^b	10° 20' 38.6478"N	77° 45' 20.7828"E	3973.097
10.	Wellington	Nilgiris	ICg10	41.10 ^e	11° 21' 44.4204"N	76° 47' 21.228"E	6085.958

150 *Mean of three replications

151 Means in a column followed by same superscript are not significantly different by Duncan's Multiple Range Test at P
 152 =0.05



153 **Fig. 1. Appearance of *C. gloeosporioides* culture on PDA medium. a) Front side, colonies**
 154 **grown were initially white-orange and then turned to greenish-grey. b) reverse side,**
 155 **colonies were brownish-orange to grey**
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157 3.2. Pathogenicity of *C. gloeosporioides*

158 The virulence of different isolates of *C. gloeosporioides* was tested in pot culture
 159 under protected cultivation. The results showed that the isolates induced the anthracnose
 160 symptoms on the inoculated anthurium leaves as like the natural symptoms. Among the ten
 161 isolates, ICg1 was found to be the most virulent isolate.
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163 **Table 2. Artificial inoculation method for pathogenicity test for anthurium anthracnose**
 164 **caused by *Colletotrichum gloeosporioides***

S.No.	Inoculation method	Per cent Disease Index(PDI)*
1.	Spraying spore suspension	36.54 ^c
2.	Injury by pinprick	47.77 ^c
3.	Pin prick + spore suspension spray	64.38 ^a
4.	Leaf clipping method	29.97 ^f
5.	Injury by carborandom powder	53.32 ^b
6.	Pin prick + mycelia disc	42.21 ^d
7.	Control (water spray)	-
CD (P=0.05)		1.170

165 *Mean of three replications

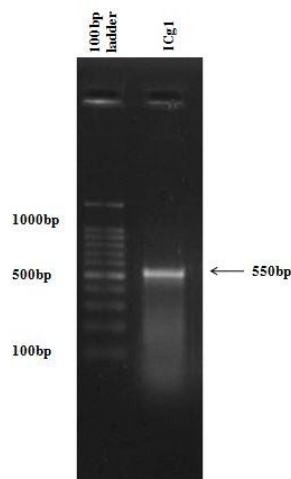
166 Means in a column followed by same superscript are not significantly different by Duncan's Multiple
 167 Range Test at P =0.05

168 The different inoculation techniques were experimented with the virulent isolate
 169 ICg1. The result showed that pinprick+spraying spore suspension method has recorded the
 170 highest disease incidence of 64.38 per cent disease index (PDI) followed by carborandom.
 171 injury by pin prick (47.77 PDI) and pin prick+mycelia disc (42.21 PDI). The least percentage
 172 was observed in leaf clipping method (29.97 PDI) (Table 2).

173 Several studies have been conducted for comparing different inoculation methods
 174 against various pathogens for screening different varieties. Co *et al.* (2008) evaluated three
 175 different smut inoculation techniques: soaking, wounding along with paste and pastes without
 176 wounding in sugarcane seedlings and found wounding along with paste method was the best
 177 method. Baayen and Schrama (1990) compared five stem inoculation methods with respect to
 178 phytoalexin accumulation and *Fusarium* wilt development in carnation and found injection
 179 method to be the more effective method.

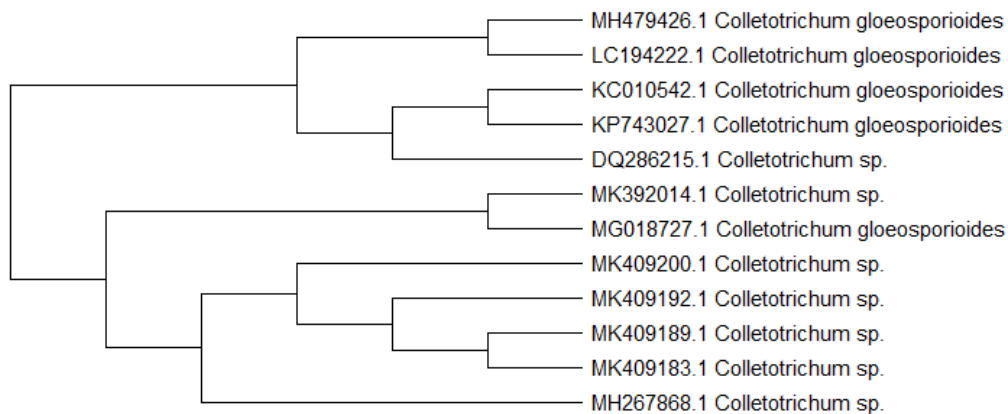
180 3.3. PCR amplification and sequence analysis

181 A PCR product of approximately 550 bp was obtained using primers ITS1 and ITS4
 182 (Fig. 2). The amplicons were purified from agarose gel and sequenced. The sequences
 183 obtained from the mycelia isolated from leaves resulted more than 92% identical to ITS
 184 sequences of *C. gloeosporioides* published in GenBank database. The obtained ITS sequence
 185 (GenBank accession number MH479426) and ten ITS sequences of the genus *Colletotrichum*
 186 retrieved from GenBank, were used to construct phylogenetic trees. The molecular
 187 identification of the ICg1 isolates as *C. gloeosporioides* was confirmed by phylogenetic
 188 analysis (Fig. 3).



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Fig. 2. PCR amplification of *C. gloeosporioides* ITS primer amplified a band of 550 bp ICg1 virulent isolate. Lane 1 is 100 bp DNA Ladder and Lane 2 is ICg1



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Fig. 3. Phylogenetic tree based on maximum parsimony analysis of the ITS sequences of *Colletotrichum gloeosporioides* (GenBank accession number MH479426) and the closest relatives from GenBank.

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The inaccuracies of identifying *C. gloeosporioides* solely by morphological criteria have been largely overcome by the use of molecular methods for differentiating *Colletotrichum* species. Ribosomal DNA sequences have been used extensively for species delineation within the genus *Colletotrichum* (Sreenivasaprasad *et al.*, 1996; Johnston and Jones, 1997), and analysis of the variable ITS regions – especially the ITS1 portion – provides sufficient information to infer phylogenetic relationships among *Colletotrichum* species (Freeman *et al.*, 2000). For this reason, ITS sequence analysis was used to verify the identity of the pathogen causing anthurium anthracnose.

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4. CONCLUSION

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Among the artificial inoculation methods, pin prick + spraying spore suspension has caused the highest percentage of disease incidence in anthurium plants. For diagnostic purposes, the sequencing method developed here showed its objective value by confirming that the strain MH479426, recently isolated from a subcutaneous infection in Brazil and identified according to its morphological features as *C. gloeosporioides* (Castro *et al.*, 2001).

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COMPETING INTERESTS

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Authors have declared that no competing interests exist

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