

Original Research Article**Characterization and control of two unknown fungal strains isolated from postharvest mango spoilage****Abstract**

Ripened mangoes are perishable sometimes as they contain large amount of water and carbon sources which make it susceptible to spoilage by different fungi. This study was therefore carried out through morphological characteristics, growth characteristics and control measure of two unknown fungal strains isolated from postharvest spoiled ‘Gopalvog’ and ‘Mollica’ mango varieties. Both the colony color of fungal strain isolated from ‘Gopalvog’ and ‘Mollica’ was initially white. Surprisingly, the colony of fungal strain from ‘Gopalvog’ became grayish brown after 72 hours. The optimum mycelial growth of fungal strain isolated from ‘Gopalvog’ was obtained at pH 8, temperature 35°C and 2% glucose concentration. The optimum pH and temperature for growth of the fungal strain isolated from ‘Mollica’ were 6 and 35 °C respectively. At 6% NaCl concentration, 100% inhibition of growth was obtained for both fungi. Growth of both the fungal strain was inhibited at 2% and 0.5-2% citric acid concentration respectively.

Keywords: Mango, postharvest spoilage, fungi, isolation, inhibition

¶ Both authors are considered as first author

Introduction

Huge amounts of fruits are cultivated across the world. The recommended quantity of fruits to be consumed by a healthy adult is 230 g/day, while the current per capita consumption of fruits is reported to be less than 160 g/day (Abdulla et al., 1981). Mango was originated from India and

24 Southeast Asia and nowadays it is one of the most important fruits cultivated in tropical
25 countries (Mukherjee, 1972). Extension of mango cultivation has been occurred to several other
26 parts of the world including Africa, the Americas and the Caribbean region (Vietmeyer, 1986).
27 Day by day consumption of mango get popularity in the developed countries(Diedhiou et al.,
28 2007). Among highest mango producing countries, Bangladesh takes 8th position in the world
29 (Amin, 2015). In Bangladesh, total production of mango is 1047849 t annum⁻¹ with an average
30 yield of 13.25 t ha⁻¹ (Amin, 2015).The potential of mango as a commercial crop is markedly
31 limited because of its high perishability, which results in considerable wastage (Mootoo, 1991).
32 Disease susceptibility due to microorganism, sensitivity to low storage temperatures and
33 perishability due to ripening and softening are serious causes of postharvest losses in mango
34 which are limiting its handling, storage and transport potential. The postharvest losses of fresh
35 mango fruits are reported to be 25 - 40% in India and 69% in Pakistan; and microbial decay
36 accounts for 17.0 - 26.9% of the total postharvest losses in Asian countries (Prabakar et al.,
37 2005). The postharvest spoilage in mangoes has been estimated to be in the range of 25-40%
38 from harvesting till they reach consumers. It is well known that mango is climacteric in nature
39 and ripen quickly after harvest. As a tropical fruit, mango is susceptible to a number of
40 physiological disorders due to low temperature during storage and even suffers from chilling
41 injury (Ding et al., 2007). At ambient temperature, harvested mango fruit at the mature stage
42 ripen quickly and have a short postharvest life, which is limited by physiological deterioration
43 related to over ripening and by pathogen development leading to decay (Johnson and Coates,
44 1993). Rapid ripening in combination with infection by microorganism is a serious cause of
45 postharvest spoilage in mango (Zheng et al., 2007). Most of cases microorganism responsible for
46 mango spoilage are fungi and bacteria where ripened mangoes are more susceptible to attack by

a variety of microorganisms (Barth et al., 2009). More than 90 fungal strains are responsible for mango spoilage (Jamalizadeh et al., 2011). “Gopalvog” and “Mollica” are the two most cultivated mango varieties in Bangladesh. These two varieties are greatly affected by postharvest spoilage. Current study was designed to characterize and control of fungi associated with the spoilage of postharvest mango varieties named Gopalvog and Mollica.

Materials and Method

Collection of fruits

Postharvest spoiled mangoes of Gopalvog and Mollica varieties were collected from Fruit Research Centre, Rajshahi, Bangladesh. The selected mangoes were separated by polyethylene bag for each type of infected fruit.

Isolation of fungi from infected fruits

The fungi responsible for the spoiled Gopalvog and Mollica mangoes were isolated on PDA (Potato Dextrose Agar) medium by following the standard procedures described by (Agostini and Timmer, 1992) with a slight modification.

Purification of culture

The fungus growing from the infected piece was removed and re inoculated on PDA medium for several times for pure culture. Single colony or sweep from the end of a hyphal tip was used as inoculum and inoculated on PDA for pure culture of respective fungus.

Microscopic observation of fungi

Mycelia from pure cultures were examined under Optika digital microscope (Italy) and was identified by comparing their morphological and cultural characteristics with previously published descriptions (Hamd et al., 2013, Barnett and Hunter, 1998)

69 **Molecular Identification of selected fungal isolates**

70 After 7 days of incubation of two fungal isolates on potato dextrose broth at $28\pm 2^{\circ}\text{C}$, DNA was
71 isolated from mycelium mat by using TIANamp Genomic DNA Kit (TIANGEN Biotech Beijing
72 co. LTD) using manufacturer's guidelines. The quality of the isolated DNA was determined
73 using 1% agarose gel electrophoresis. The primer pair ITS 1 (5'-
74 TCCGTAGGTGAACCTGCGG-3') and ITS 4 (5' -TCCTCCGCTTTATTGATATG-3') were
75 used to amplify the ITS (Internal Transcribed Spacer) region which is the universal fungal
76 primers using for identification of fungi (Hinrikson et al., 2005).

77 **Growth profiling of both fungi**

78 Potato Dextrose Agar (PDA) media was used to study the colony morphology whereas Czapeck
79 Dox Agar (CDA) and Sabouraud Dextrose Agar (SDA) media were prepared to compare the
80 morphology with PDA media. After 7 days of growth of fungi on the plates, different
81 morphological characteristics of colony such as form, elevation, margin, colour, size, surface,
82 and dry weight were observed on three different media and classified according to the cultural
83 characteristics described in (de Hoog and Guarro, 1995). Different characteristics were identified
84 for the growth profiling of the two fungal strains. Different carbohydrates such as glucose,
85 fructose, sucrose and starch were added as sole carbon source to the medium at 2% concentration
86 instead of dextrose to check the effect of them. The effect of temperature on the growth of fungi
87 was identified by incubating both the fungi at 5°C , 15°C , 25°C and 35°C at $28\pm 2^{\circ}\text{C}$ for 7 days.
88 The effect of pH on the growth of the two fungal strains was identified by inoculating both the
89 fungi into the PDA medium of pH of 6.0, 7.0, 8.0 and 9.0. Lastly, dry weight of all the fungi was
90 measured.

91 Study on cellulolytic activity

92 Cellulolytic activity of the fungi was tested using Potato Dextrose liquid medium in which
 93 sterilized 3mm filter paper was inserted as a source of cellulose. Then, 5 mm diameter plug of a
 94 7 days old colony of both fungal isolates were inoculated in the PD liquid and incubated at
 95 $28\pm 2^{\circ}\text{C}$ for 7 days and lastly flasks were observed to check the cellulose degrading ability of
 96 both fungi.

97 Control Measure by aqueous of spice and plants extract

98 Aqueous extracts of bulb of *Allium sativum*, root of *Borussus flabellifer* and leaves of *Scaparia*
 99 *dulcis*, *Pandanus odoratissimus* and *Withania somnifera* were used to investigate their
 100 effectiveness on the growth of the fungal strains.

101 Control Measure by treating with NaCl

102 The effect of salinity on the growth of the fungal strains was carried out by incubating the fungus
 103 in various NaCl concentrations- 0.5%, 1%, 2%, 4%, 6% (w/v).

104 Control measure by citric acid

105 Citric acid is one of the predominant organic acids present in mango. To observe the effect of
 106 citric acid, different citric acid concentrations of 0.25%, 0.5%, 1% and 2% (w/v) were added into
 107 the potato dextrose liquid medium and pH was adjusted to 6.5. All the inhibition percentage were
 108 measured by the following formula,

$$109 \quad \%I = \frac{C-T}{C} \times 100$$

110 Where, I= Percentage of inhibition, C= radial growth in control, T= radial growth in treatment.

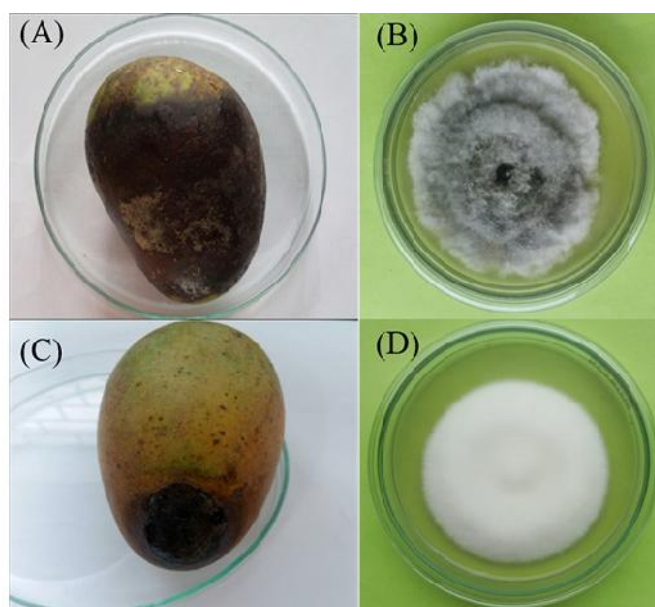
111 Statistical analysis

112 All data are the average of triplicates. All the graphs and standard error were analyzed using
113 Microsoft Excel 2016.

114 **Results**

115 **Isolation of fungi**

116 The two unknown fungal strains i.e. fungal strain-1 and fungal strain-2 were from post-harvest
117 spoilage of mangoes of Gopalvog and Mollica varieties which is showed in figure 1.

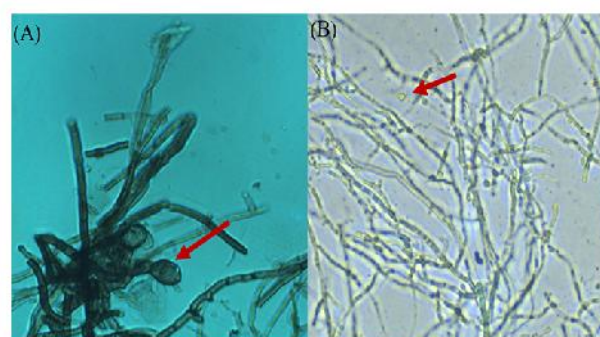


118
119 **Figure 1. Isolated strains from the postharvest spoiled mangoes.** (A) And (C) are the selected
120 postharvest spoiled mangoes. (B) and (D) are the pure culture of fungal strains named stain-1 and
121 strain-2 isolated from Gopalvog and Mollica respectively.

122 **Microscopic Identification**

123 Mycelia of the two fungi were examined and identified under microscope. In fungal strain 1,
124 colonies grew faster, mycelium was fine threaded shape and the color was white from the front
125 initially and became grayish brown in time. In addition, hyphae of the fungal strain-1 were

126 branched, broad and the diameter of hyphae was about $63.35\mu\text{m}$ and no spore was appeared. On
 127 the other hand, in fungal strain-2, colonies were very fast growing and appeared cottony to
 128 fluffy, colony color was white from the front and the shape of hyphae was very fine thin thread
 129 like. Hyphae were about $14\mu\text{m}$ in diameter and spore was not found. Microscopic view of both
 130 the fungal strain are given in figure 2.



131
 132 **Figure 2. Microscopic observation of the isolated fungal strains.** (A) Shows thicker hyphae
 133 with conidia indicated by arrow. (B) Shows thinner hyphae with spore indicated by the arrow.

134 Molecular Identification

135 DNA isolated from the fungal strains showed high molecular weight and bright band on 1%
 136 agarose gel electrophoresis where band 1 kb plus DNA ladder was used as a marker showed in
 137 Figure 3. The consensus primers ITS1 and ITS4 were used to amplify a region of the rDNA gene
 138 repeat unit. Both the isolates yielded a single band of ~ 550 bp.

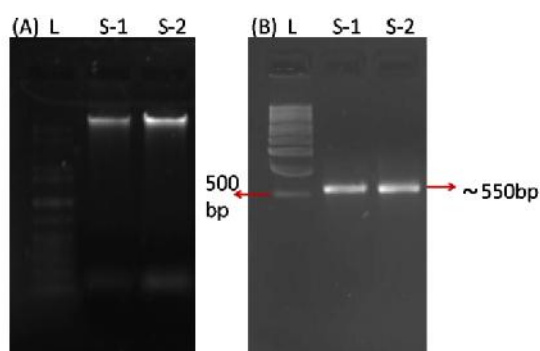


Figure 3. Molecular identification of the isolated fungal strains. L, S-1 and S-2 indicate the Ladder, Strain-1 and Strain-2 respectively. (A) High molecular weight DNA band with ladder (B) PCR amplification of ITS region showed around 550 bp band in both strain.

Colony Characterization on different media

Characterization of the colony of the two fungi were done according to (de Hoog and Guarro, 1995) by culturing them on three different types of media i.e. Potato Dextrose Agar, Czapek Dox Agar, Sabouraud Dextrose Agar. Among three types of media, SDA media increased growth of the fungal strain-1 where the growth of the fungal strain-2 was promoted by PDA media. The results are shown in table 1 & 2 and in figure 4 (A).

TABLE 1.Morphological characterization of fungal strain-1 on different growth media

Characteristics	Potato dextrose agar (PDA)	Czapek Dox Agar (CDA)	Sabouraud Dextrose agar (SDA)
1.Form	Irregular and	Irregular and	Irregular
2.Elevation	Filamentous	Filamentous	Convex
3.Margin	Raised	Cateriform	Undulated
4.Surface	Filiform	Filiform	Smooth
5.Color	Smooth	Smooth	Greyish White
6.Size (cm)	Greyish White	Greyish White	8.6cm
7. Dry weight (gm)	6.65cm 0.1464gm	5.9cm 0.1020gm	0.2845gm

155 TABLE 2.Morphological characterization of fungal strain-2 on different growth media

Characteristics	Potato dextrose agar (PDA)	Czapek Dox Agar (CDA)	Sabouraud Dextrose agar (SDA)
1.Form	Irregular and Filamentous	Irregular and Filamentous	Irregular and Filamentous
2.Elevation	Nmbonate	Nmbonate	Convex
3.Margin	Undulated	Undulated	Undulated
4.Surface	Smooth	Smooth	Smooth
5.Color	white	White	White
6.Size (cm)	4.45	2.9	4.35
7.Dry weight (gm)	0.1020gm	0.0262gm	0.0870gm

156

157 Effect of Carbohydrate on the growth of two selected fungal strains

158 To identify the effect of different carbohydrates on the growth of the two fungal strains, different
 159 carbohydrates such as glucose, fructose, sucrose and starch were added as sole carbon source to
 160 the medium at 2% concentration instead of dextrose. It was found that all the carbohydrates
 161 stimulated the growth of both isolates but glucose was more stimulatory than the other
 162 carbohydrates for the growth of fungal isolate-1 where sucrose enhanced the growth of fungal
 163 isolate-2 comparative to other carbohydrates. So, the obtained result is much closed to the
 164 referred one. The results are showed in in figure 4 (B).

165 Effect of Temperature on the growth of two selected fungal strains

166 The effect of different temperatures on the growth of both fungal strains were observed after
 167 incubation both of them at 5°C, 15°C, 30°C, and 35°C temperature for 7 days. Interestingly, both
 168 the fungal strains showed maximum mycelial growth at 35°C temperature. The results are
 169 showed in figure 4 (C).

170

Effect of pH on the growth of two selected fungal strains

pH is one of the major criteria for the optimal growth of any fungi. The mycelial growth of the two fungal strains was observed in pH values of 6.0, 7.0, 8.0 and 9.0. It was found that the fungal strain-1 showed maximum growth at pH 8.0. On the other hand, the fungal strain-2 showed maximum growth at pH 6.0. The results are showed in figure 4 (D).

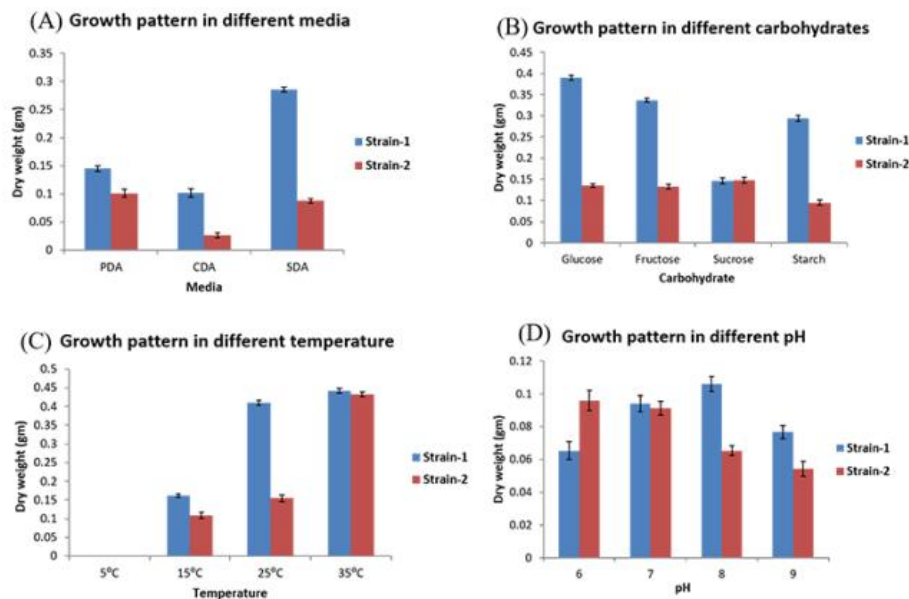


Figure 4. Growth profiling of the isolated fungal strains. (A) Highest growth was found on SDA in case of strain 1 whereas strain-1 exhibited highest growth on PDA. (B) Different growth pattern showed on different carbohydrate level. (C) Showed similar growth at 35°C temperature& (D) Optimum pH for strain-1 and strain-2 was 8 and 6 respectively.

Study of cellulolytic activity

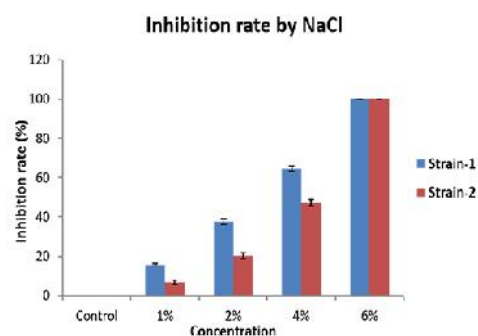
Cellulolytic activity is the ability of the cellulose enzyme to degrade cellulose. In this study, after 7 days of inoculation of fungi, it was observed that the filter papers in the cultural flasks were not degraded which indicates that both of the strains do not have any ability to degrade cellulose.

Control Measurement by treating with plant extracts

186 Different concentrations of aqueous extracts of plant parts of *Allium sativum*, *Scaparia dulcis*,
 187 *Borussus flabellifer*, *Pandanus odoratissimus* and *Withania somnifera* plants were used to
 188 investigate the inhibition rate on both fungi. In the present study, growth of both the fungi could
 189 not be controlled by 10%, 15%, 20% concentrations of aqueous extracts of the above plants.
 190 Growths of both fungi cultured with aqueous extract were close to control where the aqueous
 191 extract was absent.

192 Control Measurement by treating with NaCl

193 It was found that the increasing concentration of NaCl had a greater inhibitory effect on the
 194 growth of both fungi. It was observed that the percentage of inhibition of growth rose with the
 195 increase of the concentration of NaCl. At 6% concentration, 100% inhibition of growth of both
 196 fungal strains was observed respectively. The results are showed in figure 5.

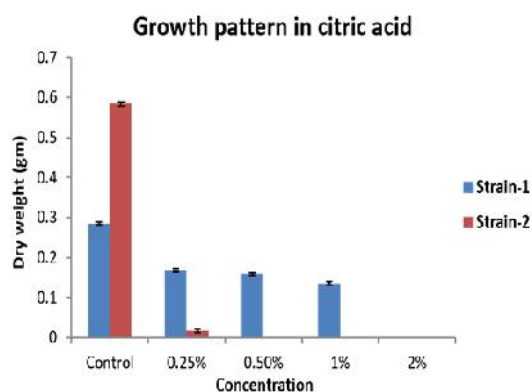


197
 198 **Figure 5. Inhibition of the fungal isolates by NaCl.** 6% NaCl showed highest inhibition for the
 199 growth of both fungal strains.

200 Control Measurement by treating with organic acid

201 To identify the effect of organic acids on the growth of both the fungi, different concentrations of
 202 citric acid e.g., 0.25%, 0.5%, 1%, 2% w/v were added to the potato dextrose liquid media. No

203 growth of the fungal strain-1 was observed at 2% concentration of citric acid. On the other hand,
 204 fungal strain-2 did not show significant growth at 0.25% concentration and no growth was
 205 noticed at 0.5-2% concentration. The results are showed in figure 6.



206
 207 **Figure 6. Growth pattern of the fungal isolates in different citric acid concentration.** 2%
 208 and 0.5-2% citric acid inhibited 100% growth of fungal strain-1 and strain-2 respectively

209 Discussion

210 Mango is one of the most popular fruits in the tropical region and its consuming rate is
 211 increasing in the developed countries day by day (Diedhiou et al., 2007). But one of the reasons
 212 for not being economically much important fruit in the world is its susceptibility to postharvest
 213 diseases (Al-Najada and Al-Suabeyl, 2014). Ripened mangoes are more susceptible to attack by
 214 a variety of microorganisms and several studies found that the main microorganisms that cause
 215 mango spoilage are fungi and bacteria (Barth et al., 2009, Jamalizadeh et al., 2011). “Gopalvog”
 216 and “Mollica” are two most cultivated varieties of mango in northern region of Bangladesh.
 217 Postharvest spoiled mango of those two varieties was collected and two unknown fungal strains
 218 i.e. fungal strain-1 and fungal strain-2 were isolated from them in PDA media. Mycelia of the
 219 two fungi were identified by comparing with the previously published descriptions in several

220 studies (Hamd et al., 2013, Barnett and Hunter, 1998) and it was found that both the colonies of
221 fungal strain 1 and 2 show different characteristics. Isolated DNA from both fungi were
222 amplified and run on gel electrophoresis which confirms the presence of the region which are
223 specific for fungi. Colony characterization of the two fungi were done according to (de Hoog and
224 Guarro, 1995) by culturing them on three different types of media where SDA and PDA media
225 increased growth of the fungal strain-1 and fungal strain-2 respectively. The results showed that
226 all the carbohydrates stimulated the growth of both isolates but glucose was more stimulatory
227 than the other carbohydrates for the growth of fungal isolate-1 where sucrose enhanced the
228 growth of fungal isolate-2 comparative to other carbohydrates which are close previous
229 studies (Devi et al., 2014, Li et al., 2014). Both the fungal strains showed maximum mycelial
230 growth at 35°C temperature which were also showed in several that fungi may grow well from
231 temperature of 25 °C to 37°C (Goyari et al., 2014, Pitt and Hocking, 2009). The mycelial growth
232 of the two fungal strains showed maximum growth at pH 8.0 and 6.0 respectively. (Goyari et al.,
233 2014, Pitt and Hocking, 2009). The isolated fungi strains are not cellulolytic as they cannot
234 produce cellulase enzyme like *Trichoderma*, *Humicola*, *Penicillium* and *Aspergillus* (Lynd et al.,
235 2002). Growth of both the fungi could not be controlled by different concentrations of aqueous
236 extracts of the part extracts of *Allium sativum*, *Scapariadulcis*, *Borussus flabellifer*, *Pandanus*
237 *odoratissimus* and *Withania somnifera* plants which all have the antifungal properties described
238 in several studies (Andriani et al., 2015, Javadian et al., 2016, Shams-Ghahfarokhi et al., 2006,
239 Singh et al.). NaCl has the ability to apply stress in the growth of fungi and it was found that the
240 increasing concentration of NaCl had a greater inhibitory effect on the growth of both fungi and
241 at 6% concentration, 100% inhibition of growth of both fungal strains was observed. Several

studies have been done on the effect of those organic acids on the growth of the fungi (Mattoo et al., 1975). No growth of both the fungal strains was observed at 2% concentration of citric acid.

Conclusion

In the present study, colony morphology of both fungi grown on different media showed different characteristics. Similar characteristics were also noticed especially in color. The maximum growth of the fungal strains was achieved at Potato dextrose agar and Sabouraud dextrose agar media respectively. The optimum temperature (35°C) and pH (8 and 6) for growth of the fungal strains were successfully identified. The most efficient carbohydrates (glucose and sucrose) for growth of the fungal strains were investigated. No cellulose degrading activity was shown by both fungi. It was identified that growth of both fungi could not be controlled by aqueous extracts of five types of plant. The control measurement of growth of the fungal strains was carried out with the treatment of NaCl. With the increase in the concentration of NaCl, the percentage of growth inhibition was increased. It was noticed that the growth of both fungal strain decreased with the increase in the concentration of organic acid. These findings will assist to prevent postharvest mango spoilage attacked by the both fungal strains. If we can interfere the conditions that increase the growth of the fungi, it is possible to prevent mango spoilage.

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