Investigation of Fungi Associated with Dried Copra and Coconut Oil

Revati R. Nalawade¹, R. R. Rathod²*, R. R. Kalaskar³, M. S. Joshi², Josiya Joy¹, Y. K. Nirgude¹, U. R. Phondekar¹ and Amruta D. Gadhave¹

¹Department of Plant Pathology, College of Agriculture, Dapoli, India
²Department of Plant Pathology, Dr. B. S. Konkan Krishi Vidyapeeth, Dapoli, India
³P.R. PotePatil College of Agriculture, Amravati, India

*Corresponding author

Abstract

Fungi associated with dried copra causing deterioration of copra and copra oil were investigated under present investigation during 2017-18. For this experiment, the infected copra samples were collected from local farmers (near Dr. Balasaheb Sawnt Konkna Krishi Vidyapeeth, Dapoli) and Regional Coconut Research Station (RCRS), Bhatye (Ratnagiri) in Konkan region of Maharashtra State, INDIA. Fungi isolated from copra samples were identified on the basis of morphological and cultural characters as Aspergillus fumigates, Chaetomium spp., Lasiodiplodia theobromae, A. oryzae and A.niger. The identifications were confirmed by Chief Mycologist, MACS- Agharkar Research Institute, (ARI) Pune.

Keywords
Aspergillus fumigates, Chaetomium spp., Lasiodiplodia theobromae, A. oryzae and A.niger

Introduction

Coconut is unique among the plantation crops grown in India as a source of food, shelter and a variety of raw materials for industrial exploitation. Coconut industry provides sustainability to a million families and livelihood to about ten million people in India. It is known that the general standard of quality and the keeping properties of the world's copra are appallingly low in comparison with the standards of every other primary agricultural produce. A variety of factors may contribute to this. There are two groups of fungi which are known to degrade copra under tropical and sub-tropical conditions.

They can be categorized as the superficial and penetrating moulds which are confined principally to the genus Aspergillus (Nathanael, 1965). India is the largest producer and exporter of coconut and copra. The copra needs to be preserved in good quality for export purpose. It is important to identify the spoilage fungi associated with the deterioration of copra and investigate possible
control measures. So an attempt was made to focus on following things:

Isolation of fungi associated with dried copra and coconut oil.

Proving pathogenicity of isolated fungi.

Materials and Methods

Diseased copra and oil samples were collected from local farmers in Dapoli (nearby DBSKKV area) and RCRS, Bhatye (Ratnagiri).

Isolation of fungi associated with dried copra and oil extracted from copra

The copra samples showing symptoms of infection were collected from local farmers. The copra was sliced into small pieces and these pieces were then surface sterilized in 0.1 per cent mercuric chloride for 2 minutes. These pieces were further washed thrice in sterilized distilled water to remove traces of mercuric chloride. Such surface sterilized pieces were aseptically transferred to sterilized Petri plates already poured with sterilized potato dextrose agar (PDA) medium. Similarly, infected oil was taken with the help of pipette and single oil drop was placed at the center of Petri plates already poured with sterilized potato dextrose agar medium without spreading.

A single bit of the fungus growing on PDA was aseptically transferred on PDA slants to obtain pure culture of the associated fungus. The slants of the pure culture were preserved in the refrigerator for future use.

Proving pathogenicity of isolated fungi

Healthy copra samples were collected from Regional Coconut Research Station, Bhatye, Dist. Ratnagiri. The copra was sliced into small pieces of about 15-20 gm weight and these pieces were then surface sterilized in 1% NaOCl for 2 minutes. These pieces were further washed thrice in sterilized distilled water to remove traces of sodium hypochlorite. The slices were respectively inoculated with the isolated fungi. This was done by dipping surface-sterilized copra slices into fungal suspension produced by homogenizing 5 discs of each fungus in 10ml sterile distilled water. These slices were placed in sterile Petri plates along with two bits of 5 mm diameter of respective culture in it. Sterile cotton plug moistened with sterile distilled water was also placed in each Petri plate containing the inoculated copra sample to provide sufficient humid conditions for growth of respective culture on copra sample. The plates were sealed with plastic tape to avoid contamination of other microorganisms.

Results and Discussion

The copra samples showing symptoms of infection were collected from local farmers and the fungi were isolated from infected portion of copra and maintained on Potato dextrose agar medium under laboratory condition for multiplication.

Colonies started developing around inoculated bits after 48 hrs of inoculation in first sample, followed by 48 hrs, 44 hrs and 38 hrs in second, third, fourth and fifth sample respectively. Five species of fungi were isolated from infected copra and coconut oil samples. These were Aspergillus fumigates, Chaetomium spp., Lasiodiplodia theobromae, A. oryzae and A.niger (Plate I and Plate II).
Plate I

D.C.1 Aspergillus fumigates

D.C.2 Chaetomium spp

D.C.3 Lasiodiplodia theobromae
Plate.II
D.C.4 Aspergillus oryzae

O.S.1 Aspergillus niger

Plate.III Proving pathogenicity of isolated fungi

A. Inoculation of isolated fungi on copra sample
B. Growth of fungi on copra sample

![Image of fungi growth](image)

**Table.1** Percentage of individual fungi isolated form diseased samples

<table>
<thead>
<tr>
<th>Fungi</th>
<th>% Frequency *</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Aspergillus niger gr.</em></td>
<td>83.33%</td>
</tr>
<tr>
<td><em>A. fumigates gr.</em></td>
<td>85%</td>
</tr>
<tr>
<td><em>A. oryzae</em></td>
<td>76.66%</td>
</tr>
<tr>
<td><em>Chaetomium spp.</em></td>
<td>50%</td>
</tr>
<tr>
<td><em>Lasiodiplodia theobromae</em></td>
<td>46.66%</td>
</tr>
</tbody>
</table>

*= Average of three replications

The identity of cultures was confirmed by Chief Mycologist, MACS- Agharkar Research Institute, (ARI) Pune. The percentages of individual fungi are shown in Table 1.

About five species of fungi namely *Aspergillus fumigates*, *Chaetomium spp.*, *Lasiodiplodia theobromae*, *A. oryzae* and *A. niger* were isolated form infected copra and copra oil samples in the percentage of 83.33%, 85%, 76.66%, 50% and 46.66% respectively. These results are in conformity with Ward and Cook (1932), Philip (1978), Kinderlerer (1984) and Morantte *et al.*, (1986) who isolated major fungi associated with deterioration of copra which were *Aspergillus niger gr.*, *A. fumigates gr.*, *A. oryzae* and *Botryodiplodia theobromae*. Venugopal S. and Chandra Mohanan R. (2006) had found *Lasiodiplodia theobromae* to be the major fungus causing fruit rot and immature nut fall in coconut.

Isolated fungi were inoculated on healthy copra samples for proving pathogenicity. The fungi started growing on copra sample 4-5 days after inoculation. Complete growth was observed on 7-10 days after inoculation. Sporulation of inoculated fungi was observed on 10-11 days after inoculation (Plate III).

**References**


How to cite this article:
doi: https://doi.org/10.20546/ijcmas.2019.810.201