Isolation and Identification of a Microorganism which Produces Non Streptomyces Pepsin Inhibitor and N-Diazoacetyl-DL-norleucine Methylester Sensitive Acid Proteases

Sawao Murao, Kôhei Oda and Yoshiyuki Matsushita

Department of Agricultural Chemistry, College of Agriculture, University of Osaka Prefecture, Sakai, Japan

Received November 22, 1972

Non pepsin inhibitor (S-PI) and diazoacetyl-DL-norleucine methylester (DAN) sensitive acid proteases producing microorganism was isolated from farm soil of Osaka Prefecture. The isolated strain was identified as Scytalidium lignicolum M-133. When it was aerobically grown on a medium consisting of glucose 5%, meat extract 1.5%, yeast extract 0.1%, KH2PO4 0.2%, MgSO4·7H2O 0.05%, at pH 3.5 and 25°C, the strain produced two acid proteases, A and B, in the culture broth.

The acid proteases A and B were not at all inactivated by S-PI and DAN. These acid proteases were expected to be a new type of acid protease from the viewpoint of the active site.

In our laboratory, pepsin inhibitor (S-PI)1,2) was isolated from Streptomyces naniwaensis, which instantaneously formed an S-PI complex with pepsin (pepsin: S-PI = 1:1) and completely inactivated the caseinolytic activity of pepsin.

The S-PI completely inactivated the caseinolytic activities of the other acid proteases produced from Rhizopus chinensis,3) Aspergillus saitoi,4) Rhodotorula glutinis K-24,5) Saccharomyces cerevisiae,6) Trametes sanguinea,7) Mucor pusillus,8) Cladosporium sp. No. 45-29) and so on, as well as pepsin.

The active site of pepsin has been studied by many investigators10–12) through the chemical modification with diazoketone compounds such as diazoacetyl-DL-norleucine methylester (DAN), and it was clarified that at the active site, there is a β-carboxyl group of an aspartic acid residue which reacts with the above reagents.

Sodek and Hofmann13,14) have isolated a peptide containing an active site residue from DAN-inactivated penicillopepsin (Penicillium janthinellum acid protease) and showed that the amino acid sequence of this peptide is quite similar to that of the active site of porcine pepsin.15) Takahashi and his colleagues16,17) studied the reaction of DAN with other microbial acid proteases from Rhizopus chinensis, Aspergillus saitoi and Mucor pusillus and with calf rennin, and assumed that the nature of the active site of acid protease was similar to each other over a very wide range of species, because these acid proteases were modified in the same manner as with pepsin.

Based on this, if there were a non S-PI and DAN sensitive acid protease, it would be expected to be a new acid protease which is different at the active site from the acid proteases reported up to date.

Therefore, as reported in the previous paper,18) the authors tried to isolate a microorganism producing a non S-PI and DAN sensitive acid protease, and obtained a strain M-133 from soil. The strain was identified as Scytalidium lignicolum M-133.

In the present paper, the isolation, identification, and cultural conditions of Scytalidium lignicolum M-133 are described.
MATERIALS AND METHODS

1. Preparation of S-PI and DAN. Pepsin inhibitor (S-PI) was prepared as reported previously.2) Diazonaetyl-DL-norleucine methylester (DAN) was synthesized according to the method described by Rajagopalan, Stein and Moore.11)

2. Assay of protease activity. Protease activity was assayed by the modified Anson method as reported previously.5)

3. Screening of non S-PI and DAN sensitive acid protease-producing microorganism. The screening of non S-PI and DAN sensitive acid protease-producing microorganism was carried out as follows. For the first screening, casein plate which contained 10 µg S-PI/ml of the medium of Table I was used. Before using, the possibility of isolating non S-PI sensitive acid protease-producing strain using S-PI casein plate was checked. That is, a cup containing 0.3 ml of pepsin (80 PU/ml), which was usually employed for the assay of penicillin, was put on the S-PI casein plate and incubated at 37°C for 18 hr. The casinolytic activity of pepsin was inactivated completely and no casinolytic ring was observed at the margin of the cup. On the casein plate without S-PI, a casinolytic ring was observed. If the acid protease is non S-PI sensitive, it must show the casinolytic ring on the S-PI plate.

Using the S-PI casein plate, streak culture was carried out in the ordinary method. As the screening source, many kinds of soil, fruit and fish were used. After 3~6 days culture at 25°C, the colony having a casinolytic ring at the margin was isolated. A second screening was performed using the isolated strains obtained, as mentioned above. The isolated strains were aerobically grown in medium II (Table I) at 25°C. Employing the supernatant of culture broth, the inhibitory activity of S-PI and DAN was assayed.

4. Assay of inhibitory activity of S-PI. A half ml of the supernatant of culture broth was incubated with 0.5 ml of S-PI solution at 37°C for 10 min and the residual activity was assayed.

5. Assay of inhibitory activity of DAN. The inhibition reaction was carried out at 15°C with DAN according to the method of Lundblad.12)

6. Identification of the strain M-133. In order to compare the morphological properties, the following type cultures were used.

(1) Scytalidium lignicolum CBS 233-57
(2) Scytalidium aurantiacum CBS 374-65
(3) Scytalidium album CBS 372-65

Color was estimated by reference to the “Guide to color standard.”19)

RESULTS AND DISCUSSION

1. Isolation of non-S-PI and DAN sensitive acid protease producing microorganism

For the screening of non S-PI and DAN sensitive acid protease-producing micro-organism, various methods such as described above were employed.

In the first screening, many fungi, five fungi imperfecti-like strains and some yeast-like strains were obtained, but most of these acid proteases were completely inactivated by 100 µg S-PI.

The authors isolated M-133 strain as the
sole non S-PI sensitive acid protease producing microorganism from farm soil collected in Osaka Prefecture University.

The acid protease of culture filtrate was not inactivated even by 100 μg of S-PI and was also not inactivated by DAN in the presence of cupric ions.

2. Taxonomical investigation of the isolated strain M–133

Taxonomical characteristics are summarized in Table II, and the photomicrographs are shown in Fig. 1, respectively.

**TABLE II. TAXONOMICAL CHARACTERISTICS OF Scytalidium lignicolum M–133**

1. Morphological characteristics
   Conidia (arthrospore): Hyaline, oblong with truncate ends, 3.5×6.5 μ×1.0–3.0 μ, produced by septation of apical part of hyphae and short branches arising at right angle.
   Chlamydospores: Brownish black, spherical to elliptical, in chains, 5.0×12.0 μ×4.5–7.5 μ.
   Hyphae: Septate, hyaline, sometimes brownish, 1.2–3.5 μ in width.
   The organ for sexual reproduction: None.

2. Cultural characteristics on various media.
   Malt extract agar: Growth abundant, flocculent, light brownish gray, reverse dark yellowish brown, soluble pigment none.
   Potato dextrose agar: Growth good, funicular, light gray, reverse radical dark olive gray, soluble pigment none.
   Czapek’s agar: Growth moderate, thin woolly, light gray, reverse dark yellow brown, soluble pigment none.

3. Physiological characteristics
   Temperature: Growing well at 25°C to 30°C, best at 30°C, growth also occurred at 37°C.
   pH: Growing well at pH 4.5 to pH 5.0, best at pH 5.0.

The isolated strain M–133 was concluded to be a species of *Scytalidium*, which belongs to a group of arthrospore-forming imperfect fungi, because it produced two kinds of spores (arthrospore and chlamydospore) under the same conditions.

In the genus *Scytalidium*, *S. lignicolum*, *S. aurantiacum*, and *S. album* are known. The morphological and physiological characteristics of M–133 were compared with their descriptions. In addition, type cultures obtained from Centraalbureau voor Schimmelcultures, The Netherlands (CBS) were cultured in parallel with M–133.

According to the description, *S. lignicolum* has an optimum temperature for growth at 29°C and grows even at 36°C, whereas the other two species have a good growth at 25°C but can not grow at 36°C. On malt extract agar, the mycelium of *S. aurantiacum* was yellow red and *S. album* produced pale yellow pigment into the medium, but these characteristics were not observed in the type culture of *S. lignicolum*.

When the type culture was cultured on malt extract agar, the growth of *S. lignicolum* was good, whereas those of the other strains were poor and the chlamydospore formation was slow. The diagnostic characteristics of type cultures were compared with those of the M–133 strain.

The M–133 strain had an optimum temperature for growth at 30°C, even grew at 37°C, the growth of the colony was good, and there was not the characteristic pigmentation at the mycelium and into the medium.

From these results, it was found that most of the morphological and physiological characteristics of M–133 resemble those of *S. lignicolum*. 

![Fig. 1. Photomicrograph of Scytalidium lignicolum M–133.](image-url)
Therefore, the authors identified the M–133 strain as *Scytalidium lignicolum*.

### 3. Culture conditions for acid protease formation

Among the carbon sources employed, glucose or mannose was best for acid protease production. Fructose and sucrose were suitable, but maltose, dextrin, glycerol were not.

The effect of several organic and inorganic nitrogen sources on acid protease production was also examined. Meat extract was the most suitable nitrogen source for acid protease production.

In addition, the effect of salts, vitamins, initial pH, culture temperature and so on was tested.

The interesting fact was that the initial pH was a very important factor for acid protease production, as previously reported. The acidic pH (pH 2.0–3.5) was suitable for acid protease production. When cultured at pH 4.5, moderate growth was observed, but acid protease production was very scant. The maximal formation of acid protease was observed when the medium in Table IV was used under the following conditions: (Temperature; 25°C, aeration; 14 liter/min, agitation; 350 rpm, 15 liter of medium per 20 liter volume fermentor).

Seed culture was previously prepared by cultivating with 100 ml of the same medium described above, in a 500 ml volume shake flask at 25°C for 2–3 days, and inoculum size was 2% (v/v).

Figure 2 illustrates the time course of acid protease formation. The proteolytic activity in the supernatant of culture medium was highest at around 80 hr cultivation, and the pH was gradually lowered from initial pH 3.5 to 2.0.

### Table III. Effect of Initial pH on the Production of Acid Protease

<table>
<thead>
<tr>
<th>Initial pH</th>
<th>Protease activity (PU/ml)</th>
<th>Growth (mg dry weight/ml broth)</th>
<th>Final pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0</td>
<td>47</td>
<td>32</td>
<td>1.9</td>
</tr>
<tr>
<td>2.8</td>
<td>59</td>
<td>33</td>
<td>2.2</td>
</tr>
<tr>
<td>3.5</td>
<td>50</td>
<td>34</td>
<td>2.2</td>
</tr>
<tr>
<td>4.5</td>
<td>12</td>
<td>23</td>
<td>5.5</td>
</tr>
<tr>
<td>6.0</td>
<td>3</td>
<td>9</td>
<td>5.5</td>
</tr>
</tbody>
</table>

### Table IV. Composition of Culture Medium

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>5 %</td>
</tr>
<tr>
<td>Meat extract</td>
<td>1.5 %</td>
</tr>
<tr>
<td>K_H3PO4</td>
<td>0.2 %</td>
</tr>
<tr>
<td>MgSO4·7H2O</td>
<td>0.05 %</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>0.1 %</td>
</tr>
</tbody>
</table>

pH 3.5

Compared with other acid protease-producing microorganisms, such as *Rhodotorula glutinis* K–24 and *Cladosporium* sp. No. 45–2, the proteolytic activity was very low.

### 4. Preliminary test for purification

The supernatant of culture medium was concentrated to one tenth of its original volume in vacuo at 40°C. The concentrated solution was dialyzed against M/100 acetate buffer (pH 4.8). The dialyzed solution was loaded on a DEAE-cellulose column, which was equilibrated with the same buffer described above, and eluted with linear gradient of NaCl. As reported previously, the authors...
found that *Scytalidium lignicolum* M–133 produced two acid proteases, A and B. The inhibition of the protease activity by S–PI and DAN was tested using crude acid proteases. Both enzymes were not inactivated by S–PI (1,000 µg) and DAN, respectively.

In the next paper, the purification of A and B acid proteases and its general properties will be reported.

REFERENCES