Characterization and Biocontrol Ability of Fusion Chitinase in *Escherichia coli* Carrying Chitinase cDNA from *Trichothecium roseum*

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Z. Naturforsch. **61c**, 397–404 (2006); received September 19/November 23, 2005

The antifungal mechanism of mycoparasitic fungi involves fungal cell wall degrading enzymes such as chitinases. *Trichothecium roseum* is an important mycoparasitic fungus with significant antifungal ability, but studies on chitinases of *T. roseum* were poor. Here, we report a novel chitinase cDNA isolated from *T. roseum* by PCR amplification based on conserved chitinase sequences. Southern blot analysis suggested that a single copy of the gene exists in the genome of *T. roseum*. The deduced open reading frame of 1,143 nucleotides encodes a protein of 380 amino acids with a calculated molecular weight of 41.6 kDa. The fusion chitinase expressed in *Escherichia coli* has been purified by single-step chromatography. It has a pI of pH 5.4 and expresses a thermal stability, but is insensitive to pH in a broad pH range. According to expectation, *E. coli* efficiently yielded a high amount of active chitinase. Remarkably, the fusion chitinase offered high antifungal activity.

**Key words:** Fusion Chitinase, Characterization and Biocontrol Ability, *Trichothecium roseum*

**Introduction**

Biological control of plant pathogens is a potential alternative to the use of chemical pesticides (Chet, 1994). The famous biological control agent *Trichoderma* is a rather specific and effective mycoparasite of various soil-borne plant pathogenic fungi in greenhouse and field conditions. So far, the extensive studies exploring the mechanisms involved in biological control were performed. Generally, the classical mycoparasitic process involves five aspects (Carsohio et al., 1994). Among them, the production of cell wall-degrading enzymes, such as chitinase is very important. Chitinase production of mycoparasites is correlated with biological control of some fungal diseases (Schirmbock et al., 1994), because chitin is a major structural component of cell wall in many plant pathogens. To date, more than ten chitinase-encoding genes have been cloned and characterized from *Trichoderma* spp. (Hayes et al., 1994; Kim et al., 2002). Furthermore, transgenic plants containing the chitinase gene (*ech42*) and transformed *Trichoderma* strains carrying multiple copies of *prb1* showed a higher resistance to several fungi (Flores et al., 1997).

*Trichothecium roseum* is another important mycoparasitic fungus and has been used as a potential agent for controlling a number of plant pathogens. The first report was on its biocontrol application against most plant pathogens causing storage rot of onions (Rod, 1984). Since then, several studies on its biological control ability were reported. For example, Urbasch (1985) and Huang and Kokko (1993), respectively, isolated *T. roseum* and tested its antifungal activity against plant pathogens such as *Pestalotia funereal* and *Sclerotinia sclerotiorum*; Kumar and Jha (2002) isolated *T. roseum* as well and designed trial controlling soybean rust by using the fungus. Exhilaratingly, they obtained a very successful biological effect. In addition, effective inhibition of plant pathogens, like *Fusarium oxysporum*, *Bipolaris oryzae*, *Alternaria* spp., *Ganoderma* spp., *Rhizoctonia solani*, *Curvularia* spp., and *Pyricularia oryzae*, has been studied for *T. roseum* as well (Balasubramanian et al., 2003). Culture filtrates of *T. roseum* have confirmed the effective inhibition of a broad spectrum of plant pathogens, though substances in the culture filtrates are very complex and unclear.

Chitinases from *T. roseum*, in comparison with *Trichoderma* spp., were poorly studied, though in-
terest has slowly increased due to the potential role as control agent for plant pathogenic fungi. The studies on characterization of purified chitinase from *T. roseum* have been reported recently (Li et al., 2004). A high level chitinase activity appeared in a broad range of pH 4.0–7.0, at 40 °C. Just as ThEn-42 (De La Cruz et al., 1992), it offered efficacious inhibition of some plant pathogens tested, particularly of *Verticillium dahliae*. However, it is difficult to develop further studies of this chitinase for two reasons: firstly, the expression of *T. roseum* was poor and inefficient, and secondly, the purification procedure was very complex and tedious. Therefore, cloning and over-expression of the chitinase gene are urgent. Here, we describe the cloning of chitinase from *T. roseum* and expression in *E. coli*. Properties and potential application of the fusion protein are evaluated.

**Materials and Methods**

**Fungus strains, plasmids, and culture conditions**

*Trichothecium roseum* s24 isolated from Shandong cotton field is part of a native strain collection held at Jilin University, Changchun/Jilin, China. In order to extract genomic DNA and total RNA, this strain was inoculated (10⁵–10⁶ conidia per ml) in medium according to Schirmbock et al. (1994) (0.68 g KH₂PO₄, 0.87 g K₂HPO₄, 0.2 g KCl, 1 g NH₄NO₃, 0.2 g MgSO₄·7H₂O, 0.2 g CaCl₂, pH 5.0) supplemented with 1 g glucose. Mycelia were grown on a rotary shaker (200 rpm) at 25 °C. After 2 d, 30 g colloidal chitin were added to the broth medium. Incubation was continued for an additional 6 d. Then, mycelia were collected by filtration, washed with sterile water, frozen in liquid nitrogen, and then stored at −80 °C. Strain JM109 of *E. coli* was transformed with recombinant plasmids and cultured in LB broth to select transformants. pET22b (+) (Novagen) was used as the cloning and expression vector.

**DNA/RNA manipulation**

Total genomic DNA from *T. roseum* was extracted followed a standard method (Hohn and Desjardins, 1992). Total RNA from *T. roseum* s24 was extracted according to the method of Jones et al. (1985) with a slight modification. In order to synthesize cDNA, poly (A) mRNA was isolated and purified by using mRNA Purification-Kit (Stratagene). DNA probes were labeled with Random Primer Labeling Kit (TaKaRa).

**Cloning of the chitinase gene cDNA from *T. roseum***

A pair of degenerated oligonucleotide primers that code for conserved amino acid sequences was designed as follows: 5′-CA(G/A)CA(G/A)CA(T/C)(C/G)(A/C)(G/T/C)(T/A)(C/T)(C/T)(A/C)CCAACCTGG-3′ (Up), 5′-CA(G/A)CA(G/T)CA(G/A)CA(C/A)(A/T)G(A/C)(G/C)(T/C)(C/T)CCC(A/G)AA(T/A/C)ACCGTTC(G/C)CCA-3′ (Dp). PCR was carried out for 35 cycles (1 min at 94 °C, 1 min at 50 °C, and 2 min at 72 °C) with total DNA fragments as templates. The PCR products were sequenced. Based on the DNA sequence and as comparison to the endochitinase gene cDNA (*ThEn-42*) from *Trichoderma* spp., A1 (5′-CCGCTGAAGATCATCCTCG-3′), S1 (5′-TACGGTGAGAAGGAGGACCA-3′) and A2 (5′-AGGGCGCTGGAGAAGCGTGTAGAC-3′), S2 (5′-TAGGGTGCACCCTTGTTAGAC-3′) were designed as two pairs of RACE primers. The two end sequences of the target gene were obtained according to the RACE method of Frohman et al. (1988), but with some modifications. First-strand cDNA fragments were synthesized according to manufacturer’s recommendations by using an improved oligo-dT11 primer [5′-CAGTGCT(G/A/C)-3′]. 5′-End Full RACE Core Set (TaKaRa) was used as follows: synthesized cDNA (hybrid DNA-RNA) was digested with RNase H (DNase free). Subsequently, circularization of single-stranded cDNA was carried out with T₄ DNA ligase. First PCR products were obtained by using the circle strand cDNA as template and A1and S1 primers. DNA amplification by nested PCR was performed with A2 and S2 as primers. Two ends of the nested PCR products were sequenced. The ORF fragments of chitinase cDNA were amplified by using RT-PCR with a pair of specific primers (OF, 5′-acgaattcgATGTTGGGTCTCC-3′; OD, 5′-gctccgagAAGTATGAAACC-3′) the italic small letters mentioned are convenient for the next DNA manipulation). The PCR products were completely digested with EcoRI and XhoI and ligated into pET22b (+) digested by using the EcoRI and XhoI too. The ligation product was electroporated in *E. coli* JM109. Transformants were selected in LB agar supplemented with ampicillin (100 µg/ml).

**Genomic Southern blot analysis of the cloned chitinase gene**

Total genomic DNA from *T. roseum* was digested to completion with the following restriction enzymes:
enzymes: BamH1, EcoRI, and XbaI, electrophoresed on 0.8% agarose gel and blotted onto Hybond N membrane (Amersham) using capillary transfer. The cloned chitinase gene fragments (ORF) above were used as probes. Southern blot was performed under high-stringency conditions.

Production and purification of chitinase

E. coli JM109 was grown at 30 °C till the culture reached an OD600 of 0.4. Incubation was continued for 4 h after adding IPTG as inducer. Cells were harvested by centrifugation, washed, and crushed with glass beads. Total proteins were dissolved in solution I [20 mm Na2HPO4 (pH 7.2), 0.5 mm NaCl]. A fraction of soluble proteins was obtained after centrifugation at 4 °C for 30 min at 14,000 × g and was loaded onto a Ni2+-NTA-agarose column (3 ml bed volume). The column was successively washed with binding buffers containing 100 mm imidazole, and then the immobilized proteins were eluted with 500 mm imidazole. His-tagged chitinase was dialysed against solution I and stored at 4 °C for further use. Protein concentrations after purification were determined by using bovine serum albumin (BSA) as the standard (Bradford, 1976). The efficiency of this purification procedure was checked by SDS-PAGE.

Chitinase activity assay

Chitinase activity was evaluated by using 4-methylumbelliferyl-β-N,N',N''-triacetylchitotriose [4-MU-(GlcNAc)3] (Sigma). A total volume of 300 µl reaction system was prepared as follows: 250 µl of reaction buffer, 30 µl of sample, and 20 µl of substrate (0.1 µg/µl) were mixed and incubated at 40 °C for 4 h; the reaction was terminated by adding 1 volume of 0.2 m Na2CO3. Chitinase activity was measured by the mean fluorescence estimated in a Turner fluorometer (model 450; 340-nm interference filter and 415-nm cut filter). One unit of chitinase activity was defined as the amount of enzyme required to release 1 µmol of 4-methylumbelliferone in 60 min. Specific activity was expressed as units per milligram of protein.

SDS-PAGE and activity staining of chitinase

Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed according to Laemmli (1970): Samples were mixed with an equal volume of 2 times Laemmli sample buffer containing 1% Triton X-100, and the mixtures were heated at 60 °C for 5 min. The gel was stained with 0.05% Coomassie brilliant blue R-250 for identification of the purified protein. In order to identify the chitinase activity, another half of the gel running in the mean time was transferred into the washing buffer [1% casein, 2 mm EDTA, 40 mm tris(hydroxymethyl) aminomethane-HCl (Tris-HCl, pH 8.0)] for the removal of SDS from the gel and the renaturation of the chitinase. After several washings in double distilled water and sodium acetate buffer (pH 4), the gel was covered with 1% low melting-point agarose supplemented with 0.2 mm 4-MU-(GlcNAc)3 in 100 mm potassium phosphate buffer (pH 6.0). Gel was inspected for fluorescent bands under ultraviolet light after incubation at 40 °C for 120 min (Tronsmo and Harman, 1993).

Effects of pH value, temperature, and metal ions on the purified fusion chitinase

The optimal pH value of the chitinase activity of the purified fusion chitinase was assayed by the following buffers: sodium lactate at pH 3.0 and 3.5; sodium acetate at pH 4.0, 4.5 and 5.5; potassium phosphate at pH 6.0, 6.5, 7.0, and 7.5; Tris-HCl at pH 8.0. The optimal temperature of the chitinase activity was assayed in microplates placed at 20 to 80 °C in potassium phosphate buffer (pH 6.0). The enzyme thermal stability was determined after preincubation in the temperature range of 20 to 80 °C for 5 h without the substrate, then the residual chitinase activity was checked under standard assay conditions. To determine the influence of metal ion on chitinase activity, Mg2+, Cu2+, Ca2+, Zn2+, Mn2+, Ba2+, Fe2+, Hg2+, Co2+, or Ag+ were tested, respectively. Each metal ion was added at a final concentration of 1 mm in potassium phosphate buffer (pH 6.0). All the reaction mixtures contained chitinase fraction and 4-MU-(GlcNAc)3 as the substrate.

Isoelectrofocusing (IEF) of the fusion chitinase

PhastGel IEF 3–9 was chosen to identify the pI of fusion chitinase. The IEF procedure was carried out according to the Phast-System User’s Manual. Standard pI marker proteins (Pharmacia) contained the following members: trypsinogen-9.3, lentil lectin basic subunit-8.65, lentil lectin middle subunit-8.45, lentil lectin acidic subunit-8.15, myoglobin basic subunit-7.35, myoglobin acidic subunit-6.85, human carbonic anhydrase B-6.55, bo-
vine carbonic anhydrase B-5.85, β-lactoglobulin A-5.20, soybean trypsin inhibitor-4.55, and amyloglucosidase-3.50.

**Antifungal activity of the purified fusion chitinase**

To check if the chitinase had antifungal activity, several economically important plant pathogens were selected: Alternaria alternata, Alternaria solani, Fusarium oxysporum, Magnaporthe grisea, Verticillium dahliae, Botrytis cinerea. In order to obtain enough spores, all pathogens were isolated from newly infected organisms and grown in potato dextrose agar (PDA). After sporulation, spores were collected, washed three times using sterile water, and then suspended in 0.1% (w/v) glucose. To examine the effects on spore germination, spores could be used directly; to examine the influence on elongation of germ tube, spores should be preincubated in normal conditions, till germ tube appeared. 50 μl of the fungal spore (germinated or not) conidial suspension were mixed with or without serial diluted amounts of purified chitinase in the wells of sterile depression slides, and incubated at 25 °C for 10 h. Influence results were investigated under a light microscope at 100× magnification.

**Results and Discussion**

A 977 bp DNA fragment was obtained from *T. roseum* by PCR amplification following the experiment procedures. Sequencing of the DNA (GenBank accession number, AY550119) showed that it shares 71% similarity with the *ech42* of *T. harzianum* (Garcia *et al*., 1994). Southern blot analysis suggested that a single copy of the gene exists in the genome of *T. roseum* (Fig. 1). So, it was reasonable to regard it as a chitinase gene fragment. Based on the amplified DNA and the cDNA sequence of *ech42*, we successfully obtained the cDNA information of two ends of target cDNA. Thus, a pair of specific primers (OF, OD) could be very easily designed, and we cloned the chitinase cDNA with standard RT-PCR method. The deduced open reading frame (ORF) of 1,143 nucleotides encodes a protein of 380 amino acids with a calculated molecular weight of 41.6 kDa (Fig. 2). Remarkably, the method we used was different from the classic RACE (Frohman *et al*., 1988). From the result, this is an efficacious and economical way to clone cDNA.

The fusion protein with more chitinase activity has been purified to SDS-PAGE homogenous (Fig. 3). The molecular weight of the chitinase is 46.2 kDa. 15.4 mg (total activity, 23,500.4 U) pure chitinase have been obtained per liter medium. IEF showed that the isoelectric point (pI) of fusion chitinase was pH 5.4, but for the chitinase produced by *T. roseum* it was pH 5.1 (Fig. 4). The purified chitinase has an optimal activity around pH 6, so it revealed the enzyme was fairly stable in a pH range of 4.0 to 7.0 (Fig. 5A). Under standard assay conditions (pH 6.0) except for temperature, the maximum activity was observed at 40 °C (Fig. 5B). After preincubation in the temperature range of 40 to 45 °C for 5 h, the fusion chitinase retained almost its full activity. Even temperature at 55 °C, it also retained 65% activity. However, the chitinase produced by *T. roseum* almost lost enzyme activity at temperatures above 45 °C (Fig. 5C). The result indicated that the fusion enzyme obtained a moderate thermal stability. Based on the similar properties of both chitinases, we think that they probably belong to the same one. The difference may be caused by the pET-22b (+) vector, if considering a redundant part of 4.6 kDa in the fusion protein.

Addition of selected metal ions to the reaction mixtures did not affect the chitinase (fusion chitinase or chitinase from *T. roseum*) activity appreciably, except Hg^{2+}. The metal ion Hg^{2+} inhibited both chitinases activity by similar percentage (Table I). This agrees with the results in related searches on inhibition of chitinases, and supports that chitinase inhibition by Hg^{2+} is a general characteristic of the chitinase group.

In our examination, spore germination and germ tube elongation of the tested fungi were significantly inhibited by the chitinase as compared with the phosphate buffer control (Table II). This
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Fig. 2. cDNA sequence (ORF) and deduced amino acid sequence of the T. roseum chitinase.

Fig. 3. SDS-PAGE of purified chitinase of T. roseum and fusion chitinase. (A) Coomassie brilliant blue R-250 staining. (B) Activity staining. Lanes: M, standard marker proteins; 1, purified chitinase from T. roseum; 2, fusion chitinase expressed in E. coli.

Fig. 4. Isoelectrofocusing of the chitinase protein. 1, Purified chitinase from T. roseum; 2, fusion chitinase expressed in E. coli; M, pI marker (the pH value of different members is indicated in Materials and Methods).

antifungal activity is consistent with what we have studied on T. roseum. Therefore, the over expressed fusion chitinase is promising to be utilized in practice.

T. roseum is a very useful fungus, not only in agriculture as a biological control agent (Balasubramanian et al., 2003), but also in medicine for the treatment of human diseases (Sesan, 1986). At the same time, it is also a notorious fungus for associations between the first trichothecene isolated from the fungus (Freeman and Morrison, 1949) and outbreaks of human diseases caused by mycotoxin
Fig. 5. The optimal pH, optimal temperature and thermal stability of the purified chitinase. The chitinase activity was detected at 40 °C for measuring the optimal pH value (A), at pH 6.0 for measuring the optimal temperature (B), or at 40 °C, pH 6.0 after the enzyme was incubated at 20 to 80 °C for 5 h with 4-MU-(GlcNAc)₃ as the substrate (C). Solid line, activity of the fusion chitinase; broken line, activity of chitinase from T. roseum.

Table I. Effects of metal ions on chitinases in potassium phosphate buffer (pH 6.0).

<table>
<thead>
<tr>
<th>Metal ion</th>
<th>Residual activity of chitinase (%)</th>
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<tr>
<td></td>
<td>Chitinase from T. roseum</td>
</tr>
<tr>
<td>Phosphate buffer</td>
<td>100</td>
</tr>
<tr>
<td>Mg²⁺</td>
<td>95</td>
</tr>
<tr>
<td>Cu²⁺</td>
<td>101</td>
</tr>
<tr>
<td>Ca²⁺</td>
<td>108</td>
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<tr>
<td>Zn²⁺</td>
<td>98</td>
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<tr>
<td>Mn²⁺</td>
<td>87</td>
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<tr>
<td>Ba²⁺</td>
<td>92</td>
</tr>
<tr>
<td>Fe²⁺</td>
<td>94</td>
</tr>
<tr>
<td>Hg²⁺</td>
<td>58</td>
</tr>
<tr>
<td>Co²⁺</td>
<td>100</td>
</tr>
<tr>
<td>Ag⁺</td>
<td>97</td>
</tr>
</tbody>
</table>

The values are means of data from two repetitions of the experiment, with three replicates per treatment.

(Joffe, 1974). In addition, Lorito et al. (1998) pointed out that a full-scale application of fungal biological control agents in commercial agriculture should be delayed because of the inconsistent results obtained by introducing these complex microorganisms into the dynamic environment. Utility of antifungal substances isolated from T. roseum is a significant aspect in biological control, but the yield by mycoparasites is relatively not enough. As a result, expression of chitinase gene from mycoparasites is a good way. Moreover, the fusion protein, with properties of a broad range of pH value and a moderate thermal stability, has a significant antifungal activity against a number of plant pathogens (data not shown). So further studies will open up at both aspects: (i) application of the fusion chitinase, (ii) in vivo analysis of biological function by disruption of the chitinase gene.

Acknowledgements

We are indebted to Dr. H. Wang and X. Men for technical assistance with IEF analyses and critical reading of the manuscript. This work was supported by a grant from the Major State Basic Research Development Program of China.
Table II. *In vitro* antifungal activity of the fusion enzyme.

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Germination</th>
<th>Antifungal activity (%)</th>
<th>Germ tube elongation</th>
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<tbody>
<tr>
<td></td>
<td>0.1 1 10 100</td>
<td>µg/ml 0.1 1 10 100</td>
<td></td>
</tr>
<tr>
<td>Alternaria alternata</td>
<td>8 23 68 92 100</td>
<td>5 34 52 87 94</td>
<td></td>
</tr>
<tr>
<td>Alternaria solani</td>
<td>17 27 56 81 93</td>
<td>24 57 83 95 100</td>
<td></td>
</tr>
<tr>
<td>Fusarium oxysporum</td>
<td>11 45 71 83 91</td>
<td>19 53 74 91 100</td>
<td></td>
</tr>
<tr>
<td>Magnaporthe grisea</td>
<td>15 38 64 73 83</td>
<td>30 67 87 93 99</td>
<td></td>
</tr>
<tr>
<td>Verticillium dahliae</td>
<td>22 56 89 100 100</td>
<td>43 73 93 100 100</td>
<td></td>
</tr>
<tr>
<td>Botrytis cinerea</td>
<td>16 34 55 68 82</td>
<td>25 35 69 84 89</td>
<td></td>
</tr>
</tbody>
</table>

The values are means of data from two repetitions of the experiment, with six replicates per treatment.

