Sequence analysis and regulation of a gene encoding a cuticle-degrading serine protease from the nematophagous fungus *Arthrobotrys oligospora*

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The nematode trapping fungus *Arthrobotrys oligospora* produces an extracellular serine protease (designated PII) that immobilizes free-living nematodes in bioassays and hydrolyses proteins of the nematode cuticle. Peptides were isolated from PII and partly sequenced. Three internal peptide sequences were used to design synthetic oligonucleotides, which allowed the subsequent isolation of the gene encoding PII from a genomic library. The deduced amino acid sequence indicated that PII is synthesized as a pre-proenzyme containing the mature enzyme, a signal sequence and a propeptide that are removed before the enzyme is secreted into the medium. The primary sequence of PII displayed a high degree of similarity with several other serine proteases of ascomycetes belonging to the subtilisin family. Northern analysis demonstrated that PII was expressed when the fungus was starved of nitrogen and carbon and that the expression was significantly stimulated by the addition to the medium of various soluble and insoluble proteins, including fragments of nematode cuticle. The levels of the mRNA as well as the proteolytic activity of PII were repressed in the presence of more easily metabolized forms of nitrogen (including ammonia, nitrate and amino acids) or glucose. The activity of the enzyme was almost completely inhibited by the peptide Phe-Val, as well as by the amino acid Phe, without a corresponding decrease in mRNA level. Notably, peptides with similar structures are known to be secreted by the host (nematode) and to stimulate the production of infection structures (traps) of the fungus.

**Keywords:** *Arthrobotrys oligospora*, cuticle-degrading serine protease, nematophagous fungi, Northern analysis, regulation of gene expression

**INTRODUCTION**

Nematophagous fungi comprise a group of mainly soil-dwelling fungi that can infect nematodes either by forming special mycelial structures (e.g. traps) or by infecting the nematodes as spores (endoparasites). A third group of nematophagous fungi are egg parasites that can penetrate the eggshell and infect the eggs of root-knot and cyst nematodes. Interest in the infection biology of these fungi has increased significantly during recent years, as nematophagous fungi have been considered for use in the biological control of plant- and animal-parasitic nematodes (Kerry, 1990; Grønvold et al., 1993). The development of new methods to control such nematodes is of major importance, because the traditional methods that are based on the use of nematicides and anthelmintic drugs are no longer effective or are associated with major environmental and health concerns. To achieve successful control of parasitic nematodes using nematophagous fungi, a detailed knowledge of the infection process is needed, virulence factors have to be identified and the factors controlling their activity have to be characterized.
The molecular background to the infection of nematodes by nematophagous fungi is, however, far from well understood. As for other parasitic and pathogenic fungi, the infection of nematodes involves a sequence of events: adhesion of the fungi to the nematode surface (cuticle), penetration of the nematode cuticle, followed by digestion and assimilation of the internal tissues of the killed nematode. By using various protease inhibitors, we have previously demonstrated that serine proteases probably play an important role during the penetration of nematodes by the nematode-trapping fungus *Artrobrotys oligospora* (Tunlid & Jansson, 1991). This is a soil-living nematophagous fungus that can grow both as a saprophyte and parasite by forming special hyphae, called traps. We have recently isolated and characterized an extracellular serine protease (PII) from *A. oligospora* that might be affected by applied serine protease inhibitors (Tunlid et al., 1994). The purified enzyme had a relatively broad substrate specificity and it was inhibited by typical serine protease inhibitors. PII is a glycoprotein with a molecular mass of approximately 35 kDa, it immobilized nematodes [*Panagrellus redivinus* L. (Goody)] in bioassays and hydrolysed proteins of the purified cuticle. The N terminus of PII was blocked, but the sequence of one internal peptide showed a high degree of sequence similarity with a region containing the active site histidine residue of the subtilisin family of serine proteases (Tunlid et al., 1994). Subtilisin-like serine proteases have also been suggested to be involved in the penetration of the eggshell of nematodes by nematophagous fungi (Lopez-Llorca, 1990; Segers et al., 1994), and recently part of a cDNA clone for such a protease was isolated and characterized (Bonants et al., 1995).

In this study, to obtain further information on the molecular structure of PII, we have cloned and characterized the corresponding gene, PII, including the 5' and 3' flanking regions. Furthermore, we have examined the expression and regulation of this protease under different growth conditions.

**METHODS**

**Peptide sequencing.** PII was purified from culture filtrates of *A. oligospora* as previously described (Tunlid et al., 1994). Purified enzyme (ca. 10 mg) was digested with *Achromobacter lyticus* protease (Wako Pure Chemical Industries Ltd) in 2 M guanidine hydrochloride as described by Rivièr et al. (1991), and with trypsin or staphylococcal V-8 protease in 0.2 M NH$_4$HCO$_3$. The peptides were separated by HPLC on a Brownlee C4 Aquapore column (2 x 30 mm) (Applied Biosystems). Pumps were System Gold (Beckman) and the detector a photo diode-array 990 (Waters, Millipore). Buffers used for reverse phase chromatography were (A) 0.1% (v/v) trifluoroacetic acid (TFA) in H$_2$O and (B) 0.1% (v/v) TFA, 10% (v/v) H$_2$O, 90% (v/v) acetonitrile. The gradient was 2-62% B over 60 min and 62-90% B over 3 min with a flow rate of 100 µl x min$^{-1}$. Elution was followed at 214 nm and fractions were collected manually. The peptides obtained were sequenced with a model 470A gas-phase sequenator (Applied Biosystems) equipped with an on-line phenylthiohydantoin (PTH) amino acid analyser (model 120, Applied Biosystems) according to the manufacturer's protocol.

**Isolation of *A. oligospora* genomic DNA.** *A. oligospora* Fres. (ATCC 24927) was grown in 1% (w/v) soya peptone on a rotary shaker (150 r.p.m.) at 20 °C for 3 d. The mycelium was filtered on a nylon mesh and high molecular mass DNA was isolated according to Wöstemeyer (1985).

**Generation of a PCR probe.** By aligning the obtained peptide sequences of PII with sequences of other proteases, it was possible to design three degenerate primers (oligonucleotides) for PCR amplification and hybridization: an upper (sense) primer located in the N terminus that included a BamHI site [5' AGGGATCCGAG(A)CAG(A)ACIGAC(T)T(AG)(CA)CTITGGG 3', 32-fold degenerate], deduced from the amino acid sequence EQTDSW in Peptide 1 (Table 1); a lower (antisense) primer in the C terminus that included an EcoRI site [5' /CGAATTCATT(C)TTA(G)TTIGGG(C)A(T)TCCCGT(A)GTT(T)ATA(G)TT 3', 64-fold degenerate], deduced from the amino acid sequence NIRGSPNK in Peptide 5 (Table 1); and an internal (sense) primer over the active site histidine residue (Tunlid et al., 1994) [5' AAT(C)GGAAT(C)GCGAT(C)TGTCGCICAT(C)GTG(T)CGGIGGIGACATTGC 3', 32-fold degenerate], deduced from the amino acid sequence DGNHGHTCAGTIA in Peptide 3 (Table 1). The upper and the lower primers were used for PCR amplification and the internal primer was used as a probe for Southern blot hybridizations of fragments obtained by PCR.

PCR was performed using a Boehringer Mannheim kit with genomic *A. oligospora* DNA as template. The amplification was initiated with a 3 min denaturation at 94 °C before addition of *Taq* polymerase, followed by 30 temperature cycles: denaturation, 94 °C for 45 s; annealing, 42 °C for 45 s; primer extension, 72 °C for 45 s. The amplification was terminated by heating at 72 °C for 10 min. PCR yielded three products (ca. 900 bp, 1100 bp and 1400 bp), as shown by gel electrophoresis in 0.8% (w/v) agarose gel. These were further analysed by Southern blot hybridization. The filter was hybridized with the 3P-labelled internal primer overnight at 45 °C and washed at moderate stringency (50 °C in 6 x SSC (1 x SSC is 0.15 M sodium chloride, 0.1 M sodium citrate) and 0.1% (w/v) SDS for 3 x 10 min). The primer hybridized exclusively to the smallest PCR product (ca. 900 bp), which was excised from an agarose gel and cloned into a pGEM-T vector according to the manufacturer's protocol (Promega). Several positive transformants were obtained; one of these was designated pGAP. Attempts to clone the PCR fragment using the restriction enzyme sites in the primers were not successful, probably due to the presence of an internal EcoRI site in the PCR fragment. Sequencing of pGAP showed that the PCR fragment had a size of 913 bp including the primers. That the cloned fragment encoded PII was verified by the presence of the sequences of the two primers used in the PCR amplification, the internal primer and of two other oligonucleotides deduced from peptide sequencing.

**Cloning of the PII chromosomal gene.** Genomic DNA from *A. oligospora* was digested with EcoRI, BamHI, HindIII and PstI (single or in pairs) to completion. The DNA was separated by gel electrophoresis in a 0.8% agarose gel and analysed by Southern blot hybridization. The filter was hybridized with the PCR probe of PII (a 3P-labelled 966 bp *Apul*–*PstI* insert of pGAP) overnight at 65 °C and washed at high stringency (65 °C in 2 x SSC and 0.1% SDS for 3 x 10 min). In the lane with DNA digested with BamHI and HindIII, the PCR probe hybridized to a single DNA band of approximately 3800 bp. A partial genomic library was constructed by recovering such DNA fragments from an agarose gel, followed by subcloning into pUC19. Approximately 5000 transformants were screened by colony
Table 1. Amino acid sequences of internal peptides from PI1 isolated from A. oligospora

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*This sequence has been published by Tunlid et al. (1994) as TNSNGN... However, further examination of the HPLC chromatograms showed that the peptide should be TDSDGN...
were from Sigma. Fragments of cuticle were prepared from the nematode *P. redivivus* by sonication and treatment with 1% SDS according to Cox et al. (1981). Before being used in the assays, the fragments were extensively washed with water and lyophilized. Soluble proteins, peptides and amino acids were sterilized by filtration (0.22 μm pore size), and collagen and cuticle fragments were autoclaved. Proteins and amino acids were added at a concentration of 1 mg ml⁻¹, unless otherwise stated. The viability of the fungus after treating the mycelium with the various polyamino acids was evaluated by transferring the mycelium incubated with the polyamino acids to an agar plate containing CMA. The fungi were incubated and the radial growth of the mycelia was observed after 1–3 d.

Protease assays and Northern blot analysis. The hydrolytic activity of PI1 was measured by using the peptide substrate N\(^{\text{N}}\)-benzoyl-L-phenylalanyl-L-valyl-L-arginine-4-nitroanilide (Bz-Phe-Val-Arg-NA) as described previously (Tunlid et al., 1994). The proteolytic activities are given in pmol-4-nitroanilide (NA) liberated ml⁻¹ min⁻¹. RNA was isolated from the mycelia using acid guanidinium thiocyanate/phenol/chloroform (Chomczynski & Sacchi, 1987). To normalize the amount of RNA used for the Northern blots, samples of the extracted RNA were initially analysed by denatured formaldehyde agarose (1%) electrophoresis and stained with ethidium bromide (Sambrook et al., 1989). The amount of RNA was evaluated by estimating the intensity of the 28S and 18S rRNA bands in an ultraviolet transilluminator. The electrophoresis experiments were repeated with approximately equal amounts of RNA and transferred to a Hybond-N+ membrane (Amersham). The filters were hybridized at 65°C in 6 x SSC overnight with a probe to isolate the entire gene from genomic DNA. To normalize the amount of RNA, filters were also hybridized with 3000 Ci mmol⁻¹ (111 TBq mmol⁻¹) using polynucleotide kinase (Sambrook et al., 1989). After digestion of PI1 with various proteases, five internal peptides were isolated and sequenced (Table 1). Sequences from three of these peptides were used to design oligonucleotide primers for subsequent PCR amplification (Peptides 1 and 5) and hybridization (Peptide 3). The obtained PCR fragment (913 bp) of PI1 was used as a probe to isolate the entire PI1 gene from genomic DNA. The isolated genomic clone (pUBH) was shown to encode PI1 by the identification of an ORF containing the deduced sequences of the five internal peptides (Fig. 1).

Southern blot analysis of genomic DNA of *A. oligospora* showed that during hybridization under a high level of stringency (filter-washed at 65°C in 2 x SSC and 0.1% SDS) the PCR gene fragment hybridized to single bands in lanes with DNA digested with restriction enzymes having no internal recognition site in the cloned gene (*BanHI*, *EcoRV* and *HindIII*) and to two bands in DNA

RESULTS

Cloning of PI1

After digestion of PI1 with various proteases, five internal peptides were isolated and sequenced (Table 1). Sequences from three of these peptides were used to design oligonucleotide primers for subsequent PCR amplification (Peptides 1 and 5) and hybridization (Peptide 3). The obtained PCR fragment (913 bp) of PI1 was used as a probe to isolate the entire PI1 gene from genomic DNA. The isolated genomic clone (pUBH) was shown to encode...
digested with EcoRI which has one internal recognition site in PII (at position 574 in Fig. 1). However, during hybridization under moderate stringency (filter-washed at 55 °C in 6 x SSC) several minor bands appeared (Fig. 2). These results indicate that there is probably only one gene encoding PII in the genome, but that related gene(s) having some similarity to PII can be found in the genome of A. oligospora.

Nucleotide sequence

The isolated clone pUBH contained the entire nucleotide sequence encoding PII as well as the 5’ and 3’ flanking regions (Fig. 1). The identified ORF begins with a translation initiation site (ATG) codon and ends at a translation termination codon (TAA) separated by 1284 bp. The sequence immediately preceding the ATG codon fits the consensus sequences of the eukaryotic translation initiation site with a CACA at position -4 to -1 (the A in ATG being +1). Several other sequences that are homologous to eukaryotic promoter elements can be identified further upstream from the ATG start codon. At -152 bp, the sequence TATAAAA indicates a potential TATA-box and a putative CAAT-box can be identified at -243 bp (Gurr et al., 1987). A putative cap-site with the sequence TCCACTCC, which is similar to the consensus cap-site sequence YYCAYYYYY (where Y = pyrimidine) (Corden et al., 1980), can be identified 36 bp downstream from the TATA-box. The ORF is interrupted by an intron of 61 bp (position 512–573) which contains the highly conservative GT and AG nucleotide pairs in the 5’ and 3’ splice sites, respectively. Finally, the sequence ATAAA at position 1756, 106 bp downstream of the termination codon, may serve as a polyadenylation site (Gurr et al., 1987).

Amino acid sequence

The ORF translates to a polypeptide of 408 amino acid residues with a calculated molecular mass of 41 949 Da, which is larger than the experimentally determined value of 35 kDa using gel filtration (Tunlid et al., 1994),
Fig. 4. Amino acid sequence comparison of PI1 with Alp from Asp. oryzae (Tatsumi et al., 1989), Alp from Asp. fumigatus (Jaton-Ogay et al., 1992), Alp from Asp. fumigatus (Jaton-Ogay et al., 1992), Alp from Asp. fumigatus (Jaton-Ogay et al., 1992), Alp from C. acremonium (Isogai et al., 1991), Alp from T. harzianum (Geremia et al., 1993), PIP from Pae. lilacinus (Bonants et al., 1995), Prl from the entomopathogen M. anisopliae (St Leger et al., 1992), Proteinase K from Tritir. album (Joshi et al., 1995) and AEP from Y. lipolytica (Davidow et al., 1987). Aspartic acid, histidine and serine residues corresponding to active site residues are marked with an asterisk below the consensus sequence. The sequence of PI1 is bold and the consensus sequence (identical residue in at least eight of the sequences) is in italics.
primary sequence of the mature protease contains four potential N-linked glycosylation sites (Asn\textsubscript{177}, Asp\textsubscript{247}, Asn\textsubscript{311} and Asn\textsubscript{294}), which follow the general rule of Asn-X-Thr/Ser, where X is any residue except perhaps aspartate, glutamic acid or proline (Mononen & Karjalainen, 1984). The proregion lacks corresponding sites for glycosylation.

The codon usage in PII shows a very marked bias, as is common in highly expressed genes in filamentous fungi (Gurr et al., 1987). As a rule, whenever possible a T or a C is preferred in the third position. Of all codons used in PII, 45% end with a C and 30% with a T, 12% with a G and 12% with an A.

**Sequence comparison**

The data bank search showed that the nucleotide and deduced amino acid sequence of PII shared extensive similarities with both fungal and bacterial members of the subtilisin family of serine proteases. The sequence identity between PII and these proteases varied between 42 and 47% (pairwise alignments of the amino acids in the proenzyme). The primary sequences of the fungal extracellular serine proteases were aligned using the program PILEUP, which by pairwise alignment creates a dendrogram showing the similarity between the sequences (Fig. 3). This analysis resulted in several clusters of proteases. However, the sequence of PII, as well as that of protease (AEP) isolated from the yeast Yarrowia lipolytica (Davidow et al., 1987), were outside these groups. This grouping of proteases was also found on inspection of the alignments by characteristic gaps and specific residues (Fig. 4). For example, the group of Proteinase K, Prl and the cuticle-degrading protease (here designated PIP) from Paecilomyces lilacinus (Bonants et al., 1995) all contain four cysteine residues in conserved positions (residues 201, 296, 357 and 437 in Fig. 4) which are not present in the other proteases. On the other hand, a high degree of similarity between the sequences was found in regions containing the active site residues aspartic acid (Asp\textsubscript{206}), histidine (His\textsubscript{148}) and serine (Ser\textsubscript{418}) (Fig. 4). The two blocks of side-chains that form the sides of the substrate-binding S\textsubscript{1} pocket in subtilisin occur in regions of high similarity and consist of Ser\textsubscript{258} Leu\textsubscript{259} Gly\textsubscript{260} and Ala\textsubscript{284} Ala\textsubscript{285} Gly\textsubscript{286} respectively, in PII. Furthermore, the highly conserved Asn\textsubscript{247} (in PII) is important in subtilisin for stabilization of the reaction intermediate formed during proteolysis (Kraut, 1977).

To further investigate whether the similarity of sequences indicated in Figs 3 and 4 reflects phylogenetic relationships between the fungi, the serine protease sequences were also analysed by the programs available within PHYLIP version 3.5c. The distant matrix methods (FITCH, KITCH and Neighbor-joining) resulted in a clustering of the proteases almost identical to that presented in Fig. 3. A slightly different clustering of the proteases was obtained using the parsimony method (PROTPARS). As with the other methods, PII and AEP were placed at a distance from a group of proteases from T. album, M. anisopliae, Paecilomyces and Beauveria bassiana (Joshi et al.,

![Fig. 5. Expression of the serine protease PII by A. oligospora incubated in LNM + 1 mg albumin ml\textsuperscript{-1} (A), LNM + 1 mg nematode cuticle ml\textsuperscript{-1} (O), or LNM only (■). Mycelia of A. oligospora were grown for 2 d in a nitrogen-limited medium before being transferred to LNM. Fragments of nematode cuticle were prepared from the nematode P. redivivus (Cox et al., 1981). (a) The proteolytic activity of extracellular PII (in the culture filtrates), as estimated by measuring the hydrolysis of the peptide substrate Bz-Phe-Val-Arg-NA. Bars indicate SD (n = 3). (b) Northern analysis of total RNA extracted from the mycelium. Ethidium-bromide-stained rRNA (top); filter hybridized with a \textsuperscript{32}P-labelled DNA fragment of PII (bottom).](image-url)

indicating that the polypeptide encodes both the mature enzyme as well as a proregion and a signal peptide. The N-terminal region of the deduced amino acid sequence resembles a typical eukaryotic signal peptide, with regard to the basic N-terminal segment, followed by a stretch of uncharged, mostly hydrophobic residues. Based on the (-3, -1) rule (von Heijne, 1983) and pairwise alignments to the subtilisin-related serine proteases Proteinase K from Tritirachium album limber (Gunkel & Gassen, 1989) and Prl from Metarhizium anisopliae (St Leger et al., 1992), we predict that the signal peptide cleavage site is located between amino acid residues 16 and 17. Furthermore, based on alignments to Proteinase K, Prl and two alkaline proteinases (Alp) from Cephalosporium acremonium (Isogai et al., 1991) and Aspergillus oryzae (Tatsumi et al., 1989) we predict that the proregion of PII is cleaved between residues 120 and 121 (Fig. 1). The predicted molecular mass of the mature protease is 28528 Da.

Biochemical experiments have previously indicated that PII is glycosylated (Tunlid et al., 1994).
Table 2. Induction of PII by various proteins, nematode cuticle and polyamino acids

Myelia of *A. oligospora* were grown for 2 d in nitrogen-limited medium before being transferred to LNM supplemented with various proteins, nematode cuticle or polyamino acids as inducing substrate. The level of PII was determined in the culture filtrates after 24 h incubation by measuring the hydrolysis of the peptide substrate Bz-Phe-Val-Arg-NA. The values are expressed as percentages of the level of PII in the medium containing LNM + albumin (mean value = 120 pmol NA ml⁻¹ min⁻¹, for three replicates, SD = 5.4). Proteins and polyamino acids were added in concentrations of 1 and 0.5 mg ml⁻¹, respectively.

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<th>Substrate</th>
<th>Relative activity (%) [mean (SD)]</th>
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<tr>
<td>LNM + albumin</td>
<td>100.0</td>
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<tr>
<td>LNM</td>
<td>5.55 (2.63)</td>
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<tr>
<td>LNM + soya peptone</td>
<td>49.7 (6.2)</td>
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<td>LNM + collagen</td>
<td>3.88 (1.75)</td>
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<tr>
<td>LNM + gelatin</td>
<td>21.5 (10.9)</td>
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<tr>
<td>LNM + nematode cuticle*</td>
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<td>LNM + polyglycin (5000–10000 Da)</td>
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<td>LNM + poly-L-glutamic acid (600 Da)</td>
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<td>32.6 (9.9)</td>
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<td>LNM + poly-L-glutamic acid (50000–100000 Da)</td>
<td>38.9 (5.52)</td>
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<tr>
<td>LNM + poly-D-glutamic acid (2000–15000 Da)</td>
<td>0.95 (0.88)</td>
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*Fragments of cuticle were prepared from the nematode *P. redivivus* (Cox et al., 1981).

1995) and from one group formed by the proteases from *Trichoderma harzianum* (Geremia et al., 1993) and *C. acremonium*. However, the proteases from *Aspergillus* were not grouped together but found at different positions in the tree (data not shown).

**Induction of PII**

Induction of PII was tested in a series of growth experiments using nitrogen-limited cultures of *A. oligospora*, because other experiments have shown that the production and secretion of extracellular proteases in fungi can be affected by the presence of intracellular nitrogen pools (Cohen & Drucker, 1977). To obtain nitrogen-starved cultures of *A. oligospora*, the fungus was grown for 2 d in LNM containing 10 g glucose l⁻¹ and 0.2, 0.02 or 0.002 g NaNO₃ l⁻¹. The mycelium was then transferred into fresh LNM/glucose medium lacking a nitrogen source and the growth of the mycelium was examined after 24 h incubation. Some growth occurred in samples collected from the LNM/glucose medium containing 0.2 or 0.02 g NaNO₃ l⁻¹, but not in those recovered from the medium containing the lowest level of NaNO₃ (0.002 g l⁻¹) (data not shown).

When such a nitrogen-starved culture of *A. oligospora* was transferred to LNM containing BSA or purified cuticle fragments from the nematode *P. redivivus* as the sole source of carbon and nitrogen (at a concentration of 1 mg ml⁻¹), the mycelium started to produce and secrete PII into the medium (Fig. 5a). Proteolytic activity of PII was also detected in the culture filtrates from mycelium incubated in LNM only, but this activity was significantly lower than in the medium containing the protein sources (5 to 20-fold lower, depending on the incubation time and protein source). The stimulation of PII production was not paralleled by a corresponding increase in mycelial biomass [the dry weight of mycelium incubated in LNM for 24 h was 2.93 mg ± 0.32 (mean ± SD) and in LNM + albumin, 3.05 mg ± 0.28]. The effect of using a medium with a higher concentration of the inducing protein was evaluated by incubating the fungus in LNM containing 10 mg albumin ml⁻¹. After 24 h incubation, the dry weight of the mycelium incubated in this medium was about three times higher than that incubated in the medium containing 1 mg albumin ml⁻¹. However, after 24 h incubation, the activity of PII was 165 pmol NA ml⁻¹ in the medium with 10 mg albumin ml⁻¹ compared to 237 pmol NA ml⁻¹ in the medium with 1 mg albumin ml⁻¹, which indicates that the higher concentration of albumin repressed the production of PII.

Therefore, in the following experiments the inducing proteins or peptides were tested at a concentration of 1 mg ml⁻¹ or lower.

By using the cloned *PII* gene as a probe, it was possible by Northern blotting to detect the PII mRNA in the mycelium incubated in LNM, LNM + albumin or LNM + cuticle after approximately 10 h incubation (Fig. 5b). The size of the PII mRNA was estimated by using commercially available RNA size markers to be approximately 1.5–1.6 kb, which was within the range of the expected size of the PII transcript (cf. Fig. 1). The variations in the level of the PII mRNA followed approximately the changes in the activity of PII; it increased with time of incubation and it was significantly
Table 3. Effects of primary and secondary nitrogen sources and glucose on the production of PI1

Myelia of *A. oligospora* were grown for 2 d in nitrogen-limited media before being transferred to LNM + albumin, and supplements of various nitrogen sources or glucose. The level of PI1 was determined in the culture filtrates after 24 h incubation by measuring the hydrolysis of the peptide substrate Bz-Phe-Val-Arg-NA. The values are expressed as percentages of the level of PI1 in the medium containing LNM + albumin (mean value = 75.7 pmol NA ml\(^{-1}\) min\(^{-1}\), for three replicates, sd = 1040). All supplements were added at 1 mg ml\(^{-1}\) except where indicated.

<table>
<thead>
<tr>
<th>Supplement</th>
<th>Relative activity (%) [mean (so)]</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH(_4)Cl (10 mg ml(^{-1}))</td>
<td>3.53 (3.19)</td>
</tr>
<tr>
<td>NH(_4)Cl (1.0 mg ml(^{-1}))</td>
<td>17.4 (14.6)</td>
</tr>
<tr>
<td>NH(_4)Cl (0.1 mg ml(^{-1}))</td>
<td>1092 (192)</td>
</tr>
<tr>
<td>NaNO(_3)</td>
<td>7.8 (2.3)</td>
</tr>
<tr>
<td>Glutamine</td>
<td>36.8 (15.3)</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>0.27 (0.46)</td>
</tr>
<tr>
<td>Valine</td>
<td>28.9 (3.06)</td>
</tr>
<tr>
<td>Arginine</td>
<td>341 (61)</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>257 (3.8)</td>
</tr>
<tr>
<td>Phenylalanine-valine</td>
<td>3.98 (3.57)</td>
</tr>
<tr>
<td>Glucose</td>
<td>50.4 (3.57)</td>
</tr>
</tbody>
</table>

higher in the mycelium incubated in medium containing albumin or nematode cuticle compared to the mycelium incubated in LNM.

The production and secretion of PI1 were stimulated by several other insoluble and soluble protein substrates including polyamino acids (Table 2). Of the polyamino acids, the highest stimulation occurred in the medium containing poly-l-glutamic acid. A range of size classes of this peptide (from 600 to 100000 Da) induced the production of PI1. In contrast, the production of PI1 was not stimulated by poly-d-glutamic acid or polyglycine. None of the used polyamino acids inhibited the growth of the mycelium of *A. oligospora* (data not shown).

Regulation of PI1

Addition of NH\(_4\)Cl at a concentration of 1.0 mg ml\(^{-1}\) or higher to LNM containing albumin repressed the production and secretion of PI1 (Table 3). No repression occurred at a concentration of 0.1 mg NH\(_4\)Cl ml\(^{-1}\). A number of other nitrogen compounds, including NaNO\(_3\) and several amino acids, also repressed the activity of PI1. Most of the amino acids tested (Glu, Val, Arg or Tyr) repressed the production of PI1 less than NH\(_4\)Cl. One notable exception was Phe, which almost completely inhibited the activity of PI1. A peptide containing Phe (Phe-Val), also strongly inhibited the activity of PI1. The production of PI1 was also partly repressed by adding glucose to the LNM/albumin medium (Table 3).

In the absence of a protein inducer (albumin), supplements of an easily metabolized carbon source (glucose, 10 mg ml\(^{-1}\)) and nitrogen source (NaNO\(_3\), 2 mg ml\(^{-1}\)) to LNM almost completely inhibited the activity of PI1 in LNM/albumin) and it was not possible to detect the transcript of PI1.

Northern blots of RNA extracted from the mycelium incubated in the media repressing the activity of PI1 showed that for several of the nitrogen sources (NH\(_4\)Cl, NaNO\(_3\), Gln and Val) as well as for glucose, a decreased activity of PI1 was paralleled by a decreased level of the PI1 transcript (Fig. 6). However, the strong inhibition of PI1 activity after adding Phe and Phe-Val was not correlated with a corresponding decrease in the level of the PI1 transcript. The level of this mRNA was similar to that observed in cultures incubated with Val.

**DISCUSSION**

The analysis of the sequence of PI1 isolated from *A. oligospora* indicated that the deduced primary structure of PI1 is similar to that of a number of other extracellular subtilisin-like serine proteases isolated from ascomycetes, including the cuticle-degrading serine proteases PIP from the entomopathogenic fungus *Pae. lilacinus* and P1 from the entomopathogen *M. anisopliae*. Several of these proteases formed groups of more closely related sequences, but the sequence of PI1 as well as that of the yeast protease AEP, were outside these groups. These relationships can be interpreted either in terms of phylogenetic distance between the species and/or as differences in the function of the enzymes. According to recent molecular phylogeny based on the analysis of rDNA, yeast (Hemiascomycetes) form an early branch within ascomycetes, separating them from filamentous ascomycetes (Taylor et al., 1994). Within the latter, rDNA gene sequences imply an early radiation of the apothecial ascomycetes (Discomycetes) from the cleistothecial (Plectomycetes) and perithecial (Pyrenomycetes) species (Gargas & Taylor, 1995). Although there are no rDNA sequences of the deuteromycete *A. oligospora* available, the teleomorph of the closely related species *Arthrobotrys superba* was recently identified as the discomycete *Orbilia fimbriata* (Pfister, 1994).
ascomycetes from which the sequences of fungal serine proteases were analysed (Fig. 3) have either been placed among Plectomycetes (M. anisopliae, Pa. lilacinus, Aspergillus sp.) or Pyrenomycetes (T. harzianum and C. aceremonium) (Taylor et al., 1994; Radford, 1993; Curran et al., 1994; Tigan-Milani et al., 1995). Thus, the similarity of the sequences of the analysed serine protease can at least partly be explained by the proposed phylogeny of the species.

Several observations indicate that the differences in sequences between PII and the cuticle-degrading serine proteases PIP and Pr1 are also paralleled by differences in biochemical properties of the enzymes. PIP, Pr1, as well as Proteinase K, are proteins with high PI values (8-9-10.3) and they show a high level of activity with chymoelastase substrates containing aliphatic and/or aromatic amino acids (Ebeling et al., 1974; Bonants et al., 1995; St Leger et al., 1987). In contrast, PII has a low PI value (4-6) and preferably hydrolyses trypsin substrates like Bz-Phe-Val-Arg-NA (Tunlid et al., 1994). Extracellular serine proteases, except PIP, have also been purified and partly characterized from the nematode egg parasites Verticillium chlamydosporium (protease designated VCP1) and Verticillium suchiasporium (P32) (Segers et al., 1994; Lopez-Llorca, 1990). The complete sequences of VCP1 and P32 have not been reported, but it was recently demonstrated that they are serologically related to Pr1, and that the N terminus of VCP1 has a sequence related to that of Pr1 (Segers et al., 1995). Furthermore, VCP1 and P32 have high PI values and VCP1 has a substrate specificity that is similar to that of Pr1 (Lopez-Llorca, 1990; Segers et al., 1994). Whether the biochemical differences between PII and PIP, VCP1 and P32 are of importance for the ability of the enzymes to degrade components of the nematode cuticle and eggs, respectively, are at present not known. In the case of Pr1, it has been suggested that the high PI value is of major importance for the hydrolytic activity and binding of the enzyme to fragments of insect cuticle (St Leger et al., 1992).

A comparison of serine protease sequences from fungi is complicated by the fact that several fungi are known to produce multiple extracellular serine proteases which might represent different functional groups of enzymes. For example, M. anisopliae produces a serine protease (Pr2) which has a similar PI value (4-4) and substrate specificity as PII (St Leger et al., 1987). Whether A. oligospora, or any other nematophagous fungi can produce both basic and acidic serine proteases with properties corresponding to Pr1 and Pr2 is not known. However, the Southern blot analysis indicated that there might exist other genes encoding subtilisin-like serine proteases in the genome of A. oligospora apart from PII. Future studies will show whether these putative genes which are clearly divergent from PII, encode extracellular serine proteases and are expressed during the degradation of the nematode cuticle.

The growth experiments indicated that the production and secretion of PII by A. oligospora are controlled by multiple regulatory circuits which include carbon and nitrogen metabolite repression as well as induction by several soluble and insoluble proteins, like fragments of nematode cuticle. Regulation of extracellular proteases by both induction and repression has been described for a number of filamentous fungi including Neurospora crassa (Drucker, 1972), the entomopathogens M. anisopliae and B. bassiana (St Leger et al., 1988; Paterson et al., 1993; Bidochka & Khachatourians, 1988) and the human pathogen Candida albicans (Lerner & Goldman, 1993). Most of these proteases can, like PII, be induced by a variety of protein substrates. However, the results from the growth experiments using polyamino acids indicate that there are some molecular features required of the peptide that can induce the production of PII. The peptides need to contain amino acids with side-chains, since polyglycine did not stimulate the production of PII, and they need to contain L-amino acids, because the L- but not the D-isomer of polyglutamic acid stimulated the production of PII. The level of stimulation was also affected by the size of the inducing peptide. It was possible to induce the expression of PII using small peptides of L-glutamic acid (600 Da, corresponding to about 4-5 residues), but the production was further stimulated in media containing higher molecular mass fractions of this polyamino acid. These results differ from those obtained in similar experiments studying the induction of an extracellular aspartyl protease in Ca. albicans (Lerner & Goldman, 1993). This protease can be induced by a number of different soluble and insoluble protein substrates, but in contrast to PII, it can be induced by D-isomers of polyamino acids (e.g. poly-D-glutamic acid) and only by peptides having a size larger than about 8 residues. Thus the results from our experiments indicate that PII is induced by a different mechanism than that suggested for the Candida peptide permease (Milewski et al., 1988), probably involving the uptake of small peptides into the cell.

The low levels of PII, produced when the mycelium of A. oligospora is starved of both carbon and nitrogen, are probably important in the nutrient-poor soil environment for regulating the production of PII during the interaction with nematodes. A basal activity of PII would allow the transformation of the insoluble cuticle to a soluble inducer that can be transported into the cell and further stimulate the expression of the PII gene. Such a mechanism has so far not been demonstrated for the induction of fungal proteases, but it has been indicated for the induction of cellulolytic enzymes in the filamentous fungi Trichoderma reesei (El-Gogary et al., 1989). Furthermore, the proteolytic activity of PII was almost completely inhibited by the peptide Phe-Val, as well as Phe, but not to the same extent by other tested amino acids including Val, Gln and Tyr. Notably, peptides containing a high proportion of non-polar and aromatic amino acids (like Phe-Val) are present in the exudate from the nematode P. redivivus, and such peptides, as well as the amino acids Phe and Val favour the formation of infection structures (traps) of A. oligospora, particularly in media with low levels of nitrogen and carbon (Nordbring-Hertz, 1973, 1977; Nordbring-Hertz & Brinck, 1974). The biological function and mechanistic basis for the differences in the
repression of PII by the tested amino acids are not known, but the results indicate that the activity of PII can be regulated by products released from the host and by compounds known to be involved in the differentiation of the infection structures of A. oligospora. Furthermore, since the levels of the PII mRNA were similar in the mycelium incubated with these amino acids, the regulation probably occurs through mechanisms acting at both the transcriptional and post-transcriptional (e.g. translational control, processing of pre-proenzyme) levels.

The decreased level of the PII mRNA in the presence of ammonia in the medium indicates that the expression of PII could be under the control of a complex regulatory circuit controlled by the product of a global regulatory gene (ara or nit-2) as in Aspergillus and Neurospora (Marzluf, 1993). Analysis of the sequence upstream from the identified ATG start codon of PII present in the pUBH clone revealed the presence of nine putative GATA core elements (in either orientation). Six of these (with the 5' end at positions -810, -808, -789, -505 and -496 in relation to the ATG start codon) have a spacing and orientation that fulfill the requirements of a strong NIT-2 binding site in N. crassa (Chiang & Marzluf, 1994).

Isolation of PII and studies on its regulation are important steps for analysing the role of cuticle-degrading proteases in the pathogenicity of A. oligospora. A long-term goal is to use such enzymes to develop new methods to control plant- and animal-parasitic nematodes. One possibility is to mass-produce the enzyme in overexpression of the corresponding gene in a heterologous system and directly apply the enzyme to the environment of the parasites. Alternatively, genes encoding cuticle-degrading enzymes can be transferred into other organisms including hosts of the parasites or micro-organisms that are common in the habitat of the parasites. Finally, by using genetic engineering, it should be possible to alter the structure of cuticle-degrading proteases and thereby obtain enzymes with modified activities and properties. To this end, it will be important to obtain further information on how these enzymes digest the nematode cuticle.

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