NtCP56, a new cysteine protease in *Nicotiana tabacum* L., involved in pollen grain development

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Abstract

Proteinases play a critical role in developmental homeostasis and in response to environmental stimuli. Our present research reports that a new cysteine protease, NtCP56, is involved in the development of pollen grains in *Nicotiana tabacum* L. The NtCP56 gene, which encodes a protein of 361 amino acid residues with a calculated molecular mass of 40 kDa, is strongly expressed in anthers. The recombinant NtCP56 showed a high activity towards casein. Kinetic analysis revealed a $K_m$ of 2.20 mg ml$^{-1}$ and $V_{max}$ of 11.07 μg ml$^{-1}$ min$^{-1}$. The recombinant NtCP56 retained more than 50% of its maximum enzymatic activity from 20°C to 60°C with an optimum Tm range of 30–50°C. The enzyme had a maximum activity at approximately pH 6.5. Suppression of the NtCP56 gene in anti-sense transgenic tobaccos resulted in the sterility of pollen grains. Our data indicated that, as a cysteine protease, NtCP56 might play an important role in pollen development.

Key words: Cysteine protease, NtCP56, pollen, tobacco.
tapetum degeneration was retarded and the middle layer cells persisted, accompanied by aborted pollen development and complete male sterility. It was also verified that the direct targets of TDR were a cysteine protease OsCP1 and its inhibitor (Li et al., 2006).

Cysteine proteases (CPs) have been reported to be the major enzymes responsible for the hydrolysis of most of the storage proteins, hordeins and glutelin, in crops (Lee et al., 2004). CPs also contribute to the regulation of programmed cell death (Solomon et al., 1999). For instance, the expression of SmCP, a S. melongena gene encoding a cysteine proteinase, coincides with several, rather than with specific, events in developmentally regulated programmed cell death in brinjal (Xu and Chye, 1999).

In this study, a new cysteine protease gene isolated from Nicotiana tabacum L. is reported. The NtCP56 gene was strongly expressed in anthers. The recombinant NtCP56 protein showed activity towards casein proteolysis in vitro. In anti-sense NtCP56 transgenic tobacco anthers, the degradation of the tapetum was retarded and pollen development was aborted. These results provide important insights into the crucial role of NtCP56 for pollen development.

Materials and methods

Molecular cloning

The GenBank database was searched using the BLAST program (http://www.ncbi.nlm.nih.gov/BLAST/) with Nicotiana tabacum cysteine protease cDNA (GenBank accession no. AY881010) as query sequence. One Nicotiana tabacum partial mRNA sequence (GenBank accession no. U57825) encoding an endopeptidase-like protein with high similarity to Nicotiana tabacum cysteine protease cDNA was identified. Based on this sequence, five primers (primer 1: 5’-TTGATGAGAGATGGAAACCCACTTCC-3’; primer 2: 5’-AGTTGAAATGCCCAGCAACTTCC-3’; primer 3: 5’-TGTATGCTACGGAGGATAG-3’; primer 4: 5’-CTGATCAGAGGTACCCGATCC-3’; and primer 5: 5’-TTTAAACATGAAGAATTATTACTAC-3’) were designed to amplify Nicotiana tabacum NtCP56 cDNA.

Total RNA from anthers of Nicotiana tabacum L., isolated using the Trizol reagent, was used for a reverse-transcription polymerase chain reaction (RT-PCR). The first strand cDNA was synthesized in a volume of 25 μl containing about 2 μg RNA, 4 μl reaction buffer, 1 μl of 10 mM dNTP, 0.5 μl RNase inhibitor, 1 μl adaptor primer (5’-GTGATCAGAGGTACCCGATCC-3’), and 1 μl AMV reverse transcriptase. The transcription reaction was performed at 42 °C for 1 h and terminated at 65 °C for 10 min (Li et al., 2002).

A specific cDNA fragment was first amplified by using primers 1 and 2. To isolate the complete region of this gene, a 3’ race PCR was carried out by using primer 3 and primer 4 according to the instructions of the manufacturer (Takara 3’ full race kit, Japan). The PCR was carried out as follows: 94 °C for 5 min, followed by 25 cycles of 94 °C for 1 min, 56 °C for 1 min, and 72 °C for 2 min. To verify the integrity of the cDNA, sequences of the gene RT-PCR were carried out to amplify the full-length cDNA using primers 4 and 5. The PCR thermal cycles were carried out as follows: 94 °C for 5 min, followed by 25 cycles of 94 °C for 1 min, 45 °C for 1 min, and 72 °C for 2 min. The PCR product was cloned into the pMD18-T vector (Takara, Japan) and sequenced using the chain-terminating method with an ABI automated sequencer (model no. 377). The cDNA from Nicotiana tabacum L. was termed NtCP56 (GenBank accession no. EU429306).

Phylogenetic analysis

A phylogenetic tree between NtCP56 and the other major classes of plant CPs was constructed with MEGA v.2.1 by using the Neighbor-Joining (NJ) method with bootstrapping 500 replications.

Northern blotting analysis

For the RNA gel blot analysis, 25 μg of RNA was fractionated by gel electrophoresis in a 1.2% formaldehyde agarose gel, then transferred to a Hybond-N+ membrane (Amersham, UK) after UV cross-linking (Yu et al., 2003). The gene-specific DNA probe was prepared by amplifying a 356 bp 5’ cDNA fragment of NtCP56. The probe was labelled using a DIG-primer labelling kit according to the instructions of the manufacturer (Roche, Germany). Membrane-bound DNA was prehybridized for 1 h and the hybridization of probes took place overnight at 42 °C. Two subsequent stringency washes were performed at 25 °C for 10 min and at 65 °C for 30 min. The first washing solution contained 0.1% SDS (w/v), 2× SSC, and the second 0.1% SDS (w/v), 0.5× SSC. Kodak X-Omat AR-5 films were exposed to the blots (Amersham, UK).

Expression, purification, renaturation, and sequencing of recombinant NtCP56

The open reading frame minus the first 60 bp of the cDNA encoding NtCP56 was amplified. The PCR product was digested with KpnI and SacI and subcloned into the plasmid expression vector pET30a (Novagen Inc., Madison, WI, USA). Fresh colonies of BL21 (DE3) bacteria transformed with plasmid pET30a/NtCP56 were incubated overnight at 37 °C. Overnight cultures were diluted to 1:100 in a 1.0 l fermenter and grown to mid-log phase. 0.4 mM IPTG was added and cultures were incubated for 3–5 h following induction. Bacterial pellets were harvested and stored at −70 °C (Smith and Gottesman, 1989).

The resuspended bacteria were disrupted by cold sonication in a Heat Systems-Ultrasonic sonifier cell disruptor at 200 W (120×6 s) after suspension in the buffer; the insoluble protein fractions were isolated by centrifugation for 10 min at 12 000 g. The pellets were washed twice in the same buffer. The insoluble protein pellets were resuspended for 60 min.
using 3 ml g⁻¹ bacteria of buffer A (10 mM TRIS-HCl, 8 M urea, 50 mM NaH₂PO₄ and the pH adjusted to 8.0 using NaOH) and incubated at 28 °C for 1 h.

The resuspended solution were added to the Ni²⁺-NTA resin (4 ml) which was equilibrated by buffer A in a 15 ml conical tube and gently shaken at 4 °C for 1 h. Then the bottom cap was removed and the flow-through was collected. The column was washed with 20 column volumes of washing buffer B (10 mM TRIS-HCl, 8 M urea, 50 mM NaH₂PO₄ and the pH adjusted to 6.3 using HCl) and buffer C (10 mM TRIS-HCl, 8 M urea, 50 mM NaH₂PO₄ with the pH adjusted to 5.9 using HCl). The over-expressed protein was then eluted in 4 ml fractions in buffer D (10 mM TRIS-HCl, 8 M urea) (Handbook for high-level expression and purification of 6xHis-tagged proteins, Novagen).

Denatured proteins in a dialysis bag were immersed into a 50-fold volume renaturation buffer (50 mM potassium phosphate, pH 10.7, 5 mM EDTA, 1 mM reduced glutathione, 0.1 mM oxidized glutathione, 6 M/4 M/2 M/0 M urea) (Smith and Gottesman, 1989) and stirred at 4 °C for 3–4 h. Then the buffer was refreshed in turn by a reduction of urea.

Recombinant NtCP56 was processed to a mature form at pH 3.0 with cysteine (Troen et al., 1988). NH₂-terminal amino acid sequencing was accomplished by the 2,4-dinitrofluorobenzene (DNFB) method.

**Enzyme assays and kinetic studies**

Enzyme activity of purified recombinant NtCP56 was routinely assayed with casein (Sigma, USA) as substrate, using the method described by Sasaki et al. (1977) with minor modifications. Assays with up to 1% casein were carried out at 37 °C and pH 7.2, following 30 s activation incubation at 37 °C, pH 3.0, as described by previous reports (Troen et al., 1988). Casein (1 ml of 0.04–1% solution in a phosphate-buffered saline containing 6 M cysteine) was mixed with recombinant processed protein and the tubes were incubated at 37 °C for 3 min. After
incubation, the proteins were precipitated by the addition of 2 ml of a 20% tricarboxylic acid buffer and centrifuged at 12,000 g for 10 min. The absorbance of the clear supernatant was measured by a spectrophotometer at 275 nm. Controls, such as substrate and enzyme blanks, were included to obtain a background absorbance that was subtracted from the enzyme-containing tubes to yield a specific degradation of casein. The following two buffers were used: 100 mM sodium acetate (pH 3.0–5.0) and 100 mM sodium phosphate (pH 6.0–8.0). Kinetic constants were analysed by Matlab software. Leupeptin (100 μM), antipain (100 μM), and tosyl-L-lysine chloromethyl ketone (TLCK) (100 μM) were used as inhibitors.

**Anti-sense plasmid construction and generation of transgenic plants**

The 356 bp fragment of NtCP56 near the 5’ UTR was amplified and a BamHI and XbaI restriction enzyme site was created by PCR. After digestion with BamHI and XbaI restriction enzymes, the fragment was cloned into pBI121 (Clontech, USA) in an anti-sense orientation to create the 35S-anti-NtCP56 fusion construct. The fusion construct plasmid was first introduced into Agrobacterium tumefaciens LBA4404 and verified by using PCR. The empty vector (pBI121) was used as a control. Tobacco plant transformation from leaf explants was carried out as described by Lu et al. (2003).

**Histological study**

Anthers of control and transgenic tobacco were collected in a series of development stages. Anthers were fixed with FAA (5% formalin, 5% acetic acid, and 70% ethanol, by vol.) for 24 h under negative pressure and dehydrated in a set of increasing solutions (85%, 95%, and 100% by vol.) for 24 h under negative pressure and dehydrated in 50% ethanol (v/v) and dimethylbenzene and finally embedded in paraffin. Paraffin-embedded anthers were cut to a thickness of 10 μm and stained with safranine and a fast green solution.

**Results**

**Isolation and characterization of NtCP56**

A 1289 bp cDNA was cloned from *Nicotiana tabacum* L. anthers. The cDNA contained a 1083 bp open reading frame (ORF), 7 bp of 5’ untranslated region (UTR), and a 199 bp 3’ UTR. The ORF encodes a peptide of 361 residues which has a predicted molecular mass of 43 kDa (Fig. 1). This cDNA shows 36–83% similarity with other KDEL-tailed plant cysteine proteinases, including the *Solanum lycopersicum* KDEL-tailed cysteine endopeptidase, *Ricinus communis* Cys-EP (Schmid et al., 1998), *Phaseolus vulgaris* EP-C1 (Tanaka et al., 1991), *Glycine max* CysP1 (Ling et al., 2003), and *Vigna mungo* sulphidyl endopeptidase SH-EP (Akasofu et al., 1989). Then the cDNA is named as NtCP56. A phylogenetic analysis was performed using the MEGA v.2.1 programme (Fig. 2). The NtCP56 protein is located in the group which contains KDEL-tailed family members. Given this analysis, the NtCP56 protein can be classified as belonging to the subfamily C1A of papain-like cysteine proteinases (MEROPS peptidase database, http://merops.sanger.ac.uk; Rawlings et al., 2008).

The expression of the NtCP56 gene in different tissues of tobacco is determined using Northern blotting (Fig. 3).
The result showed that the NtCP56 gene is strongly expressed in anthers and weakly expressed in petal, pistil, stem, and leaf.

Expression, purification, and sequencing of recombinant NtCP56 protein

The plasmid which encodes the full-length NtCP56 minus the signal peptide was efficiently expressed in E. coli. Three hours after induction with IPTG, recombinant NtCP56 represented approximately 20% of the total bacterial protein. The molecular weight of the purified recombinant NtCP56 was estimated to be approximately 43 kDa according to the SDS-PAGE analysis (Fig. 4, Lane 3). This 43 kDa protein did not have any determinable activity (data not shown). After renaturation, the purified protein could be processed into a 33 kDa mature form at pH 3.0 (Fig. 4, Lane 4). The NH$_2$-terminal amino acid sequence of processed recombinant NtCP56 was HEDSVPP which, with four additional NH$_2$-terminal amino acids (HEDS), was compared with the predicted autocatalysis.

Biochemical characterization of processed recombinant NtCP56

The steady-state kinetics of the enzyme was studied in assays with various concentrations of casein as substrates. Processed recombinant NtCP56 obeyed the Michaelis–Menten-type of kinetics towards casein. The $K_m$ (2.20±0.04 mg ml$^{-1}$) and $V_{max}$ (11.07±0.23 μg ml$^{-1}$ min$^{-1}$) values for the recombinant enzyme are comparable to those reported for papain-like proteins from different species (Table I). The activity of the recombinant NtCP56 is pH-dependent. The highest activity was observed at pH 6.5, close to the optimal pH 7.0 of commercial papain (Hoover and Kokes, 1946). The enzyme still showed 53% of its maximum activity between pH 3.5 to pH 7.5, but only 28% at pH 8.5 (Fig. 5A), which was similar to the results of a previous report (Smith and Gottesman, 1989).

The optimum temperature for NtCP56 activity was at 30 °C. The processed recombinant NtCP56 retained more than 50% of its maximum enzymatic activity between 20–60 °C (Fig. 5B).

| Table 1. Enzymatic properties of recombinant NtCP56 |
|-----------------------------------------------------|
| **Recombinant NtCP56** | **Papain$^a$** | **Fruit bromelain$^b$** |
| $K_m$ (mg ml$^{-1}$) | 2.20±0.04 | 1.4709 | 0.7 |
| $V_{max}$ (μg ml$^{-1}$ min$^{-1}$) | 11.07±0.23 | 5.5252 | 134.2 |
| $K_{cat}$ (min$^{-1}$) | 1.025 | | |
| $K_{cat}/K_m$ (ml mg$^{-1}$ min$^{-1}$) | 0.46 | | |
| Inhibitor action (%) | | | |
| None | | 45.76% | |
| Leupetin (100 μM) | | | |
| Antipain (100 μM) | | 18.60% | |
| Tosyl-L-lysine chloromethyl ketone (TLCK) (100 μM) | | 11.02% | |

$^a$ Xiao (2005). $^b$ Zhao et al. (1999).

Phenotypic analyses of anti-NtCP56 transgenic tobacco

In order to discover the function of NtCP56 in anther development, anti-NtCP56 transgenic tobaccos were obtained. In order to ascertain whether or not the NtCP56 transgenic tobaccos...
gene plays a crucial role in the growth and development of tobacco plants, the root length and plant height of transgenic and control tobacco plants were measured each week. All 14 transgenic plants are phenotypically indistinguishable from the control plants under normal growing conditions. Compared with the control, the mRNA level of NtCP56 was reduced by 45–67% (Fig. 6A, B).

Anther development was concisely divided into six stages based on a previous classification of anther development (Feng et al., 2001). During the microspore mother cell (MMC) stage, there was no detectable morphological difference in anthers between the control and anti-sense transgenic plants. Normal microsporocytes, an epidermis, an endothecium, a middle layer, and a tapetum were found both in controls and in transformation anthers (Fig. 7A, B). Up to the tetrad stage, there was still no obvious difference in anther cellular morphology between the control and anti-sense transformations. During this period, both in the control and in transgenic anthers, microsporocytes underwent meiosis to form tetrads of four haploid microspores. The tapetal cells had begun to differentiate and their cytoplasm became deeply stained (Fig. 7C, D) (Li et al., 2006). Subsequently, the transgenic anthers had detectable morphological abnormalities. At the young microspore stage, in control anthers, microspores were released from tetrads and dispersed independently in locules (Fig. 7E). Partial tapetal cells were degenerated and the tapetum became thin (Fig. 7M). By contrast, in transformation anthers the tapetum remained relatively thick (Fig. 7N). Next, from the young pollen stage to the pollen mitosis stage, in control anthers, the uninucleate pollen developed to dianucleate pollen through mitotic divisions and dispersed in locules (Fig. 7G, I). Meanwhile, almost all tapetal cells had differentiated and degenerated (Fig. 7O). However, in the anti-NtCP56 transgenic anthers, abnormal microspores collapsed and remained undeveloped following the release from tetrads (Fig. 7H, J). Partial tapetum was still retained during this period (Fig. 7P). At the mature pollen stage, control pollen grains were full of starch, lipids, and nutrients (Fig. 7K) and the tapetum was fully degenerated (Fig. 7Q). By contrast, in transformation anthers, many pollen grains had become abnormal and did not mature (Fig. 7L, R). Large numbers of stagnated, deformed, and aggregated pollen grains were observed, and particularly in the corners of the locules (Fig. 7T). Most of the pollen grains did not stain densely, which suggested immaturity. In addition, from the young pollen stage, the cells of the endothecium gradually became fibrotic in control anthers. By contrast, in the transformation of anti-NtCP56, the endothecium cells remained loosely arrayed and did not have clear fibrotic cell walls.

In order to validate the sterility of transformation pollen grains further, the germination rate of pollen grains in vitro were determined. The germination rate of pollen grains is 41.3% of transgenic lines (102 in 247) and 5.3% of control

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**Fig. 5.** Effect of pH and Tm on the enzyme activity. (A) pH stability of processed recombinant NtCP56. (B) Thermal stability of processed recombinant NtCP56.

**Fig. 6.** Relative expression level of NtCP56 mRNA in control, anti-NtCP56-1 and anti-NtCP56-2 tobaccos. (A) Results of the semi-quantity RT-PCR by 1% agarose. (B) Quantification of the result of the semi-quantity RT-PCR.
The germination rate of control lines is about eight times higher than that of transgenic lines. This result showed most transgenic pollen grains were abnormal in development. These results suggested that the suppression of NtCP56 resulted in defects in pollen grain development.

Discussion

In this study, a gene from the tobacco anther, which is coded for a cysteine protease named NtCP56 has been isolated and characterized. Given the alignment to other plant papain-like cysteine proteases, the sequence of NtCP56 was identified as a signal peptide that would be processed prior to enzyme activation. Like other papain-like cysteine proteinases, three highly conserved catalytic residues Cys150, His286, and Asn307 constitute the catalytic triad of cysteine proteases (Guerrero et al., 1998). Another conserved residue, Gln144, is involved in maintaining an active enzyme conformation (Kamphuis et al., 1985). NtCP56 protein also contains a conserved non-contiguous ERFNIN motif (EX3RX3FX2NX3I/VX3N) which is typical for cysteine proteinases in the cathepsin L and H-like proteinases but not in cathepsin B-like proteinases (Wiederanders, 2003). A GCNGG motif was identified in NtCP56. With the exception of the central Asn (N) residue, this GCNGG motif is invariant in all ERFNIN proteinases.
and also in the cathepsin B-like proteinases (Beyene et al., 2006). In addition, a C-terminal tetrapeptide KDEL is present. It has been suggested that this is an endoplasmic reticulum (ER) retention signal (Fig. 1) (Guerrero et al., 1998).

Foreign proteins overexpressed in E. coli frequently formed insoluble aggregates or inclusion bodies in the bacterial cytoplasm (Marston, 1986). When plasmid pET30a-NtCP56 was expressed in E. coli, the recombinant proteins quickly formed inclusion bodies and could only be solubilized in strong detergents such as 8 M urea. Denatured soluble protein was eluted at pH 4.5 after purification by Ni-NTA resin. Renaturation was carried out at neutral or alkaline pH to ensure recovery of the pro-enzyme in order not to be autocatalysed (Gal and Gottesman, 1986). This might be a result of the difference between the maturation conditions in vitro and in the plant (Vernet, 1990).

The NtCP56 gene is strongly expressed in anthers. This result is analogous to OsCP1 reported in 2004, where its promoter was highly active in the anther, but not in other flower organs and vegetative organs of rice (Lee, 2004). This is different with most homologous CPs which have been reported and participated in processing and degradation of seed storage proteins (Shimada et al., 1994; Toyooka et al., 2000), fruit ripening (Alonso and Granell, 1995) and also in legume nodule development (Naito et al., 2000). But it was similar to OsCP1 isolated from rice, which is involved in pollen development (Lee et al., 2004). It has been suggested that cysteine proteinases have separate roles in different tissues (Lee et al., 2004). RD21 (Yamada et al., 2001), a cysteine protease that belongs to the papain family in Arabidopsis, has a role in the degradation of cellular proteins during leaf senescence, rather than in the degradation of seed storage protein after seed germination. Thus, it is speculated that NtCP56 is a novel cysteine protease which might be involved in anther development.

It has been demonstrated that the protease activity increased during anther development (DeGuzman and Riggs, 2000). This enhanced activity can be correlated with morphological and biochemical events during late microsporogenesis which require proteolytic enzymes. Pollen development is a dynamic process that involves protein synthesis required for the development (Lee et al., 2000). This enhanced activity can be correlated with degradation of cellular proteins during leaf senescence, rather than in the degradation of seed storage protein after seed germination. Thus, it is speculated that NtCP56 is a novel cysteine protease which might be involved in anther development.

An anti-sense approach was used to analyse the role of NtCP56 in pollen development. Until the tetrad stage, the anther development of transgenic plants was normal. However, after that, development of both tapetum and microspores was disrupted in anti-NtCP56 anthers. It suggested that NtCP56, as a cysteine protease, may participate in new protein synthesis for pollen development or aid in the degenerative process of the tapetum. It is either activating hydrolytic enzymes that degrade cellular macromolecules, or by functioning as a hydrolytic enzyme, or both (Koltunow et al., 1990). At present, the mechanism of NtCP56 during pollen development remains to be explained.

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