<table>
<thead>
<tr>
<th><strong>Title</strong></th>
<th>Brassica juncea chitinase BjCHI1 inhibits growth of fungal phytopathogens and agglutinates Gram-negative bacteria</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Author(s)</strong></td>
<td>Guan, Y; Ramalingam, S; Nagegowda, D; Taylor, PWJ; Chye, ML</td>
</tr>
<tr>
<td><strong>Citation</strong></td>
<td>Journal Of Experimental Botany, 2008, v. 59 n. 12, p. 3475-3484</td>
</tr>
<tr>
<td><strong>Issued Date</strong></td>
<td>2008</td>
</tr>
<tr>
<td><strong>URL</strong></td>
<td><a href="http://hdl.handle.net/10722/60690">http://hdl.handle.net/10722/60690</a></td>
</tr>
<tr>
<td><strong>Rights</strong></td>
<td>This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License.; Journal of Experimental Botany. Copyright © Oxford University Press.</td>
</tr>
</tbody>
</table>
RESEARCH PAPER

**Brassica juncea** chitinase BjCHI1 inhibits growth of fungal phytopathogens and agglutinates Gram-negative bacteria

Yuanfang Guan\(^1\), Sathishkumar Ramalingam\(^1,*,^+\), Dinesh Nagegowda\(^1\), Paul W. J. Taylor\(^2\) and Mee-Len Chye\(^1,†\)

\(^1\) School of Biological Sciences, The University of Hong Kong, Pokfulam Road, Hong Kong, China
\(^2\) BioMarka, School of Agriculture and Food Systems, The University of Melbourne, Victoria 3010, Australia

Received 17 May 2008; Revised 2 July 2008; Accepted 3 July 2008

**Abstract**

*Brassica juncea* BjCHI1 is a plant chitinase with two chitin-binding domains. Its expression, induced in response to wounding, methyl jasmonate treatment, *Aspergillus niger* infection, and caterpillar *Pieris rapae* feeding, suggests that it plays a role in defence. In this study, to investigate the potential of using BjCHI1 in agriculture, *Pichia*-expressed BjCHI1 and its deletion derivatives that lack one or both chitin-binding domains were tested against phytopathogenic fungi and bacteria. Transplastomic tobacco expressing BjCHI1 was also generated and its extracts assessed. In radial growth-inhibition assays, BjCHI1 and its derivative with one chitin-binding domain showed anti-fungal activities against phytopathogens, *Colletotrichum truncatum*, *C. acutatum*, *Botrytis cinerea*, and *Ascochyta rabiei*. BjCHI1 also inhibited spore germination of *C. truncatum*. Furthermore, BjCHI1, but not its derivatives lacking one or both domains, inhibited the growth of Gram-negative bacteria (*Escherichia coli*, *Ralstonia solanacearum*, *Pseudomonas aeruginosa*) more effectively than Gram-positive bacteria (*Micrococcus luteus* and *Bacillus megaterium*), indicating that the duplicated chitin-binding domain, uncommon in chitinases, is essential for bacterial agglutination. Galactose, glucose, and lactose relieved agglutination, suggesting that BjCHI1 interacts with the carbohydrate components of the Gram-negative bacterial cell wall. Retention of chitinase and bacterial agglutination activities in transplastomic tobacco extracts implicates that BjCHI1 is potentially useful against both fungal and bacterial phytopathogens in agriculture.

Key words: Bacterial agglutination, chitin-binding domain, chloroplast transformation, Indian mustard, lectin, phytopathogens, *Pichia*-expressed proteins, transplastomic tobacco.

**Introduction**

Many classes of plant lectins have been identified and they have been implicated in plant defence (Peumans and Van Damme, 1995). One well-characterized group structurally consists of chitin-binding domains, of which one such domain occurs in the mature form of hevein (Van Parijs *et al.*, 1991; Beintema, 1994). Hevein-like chitin-binding domains have also been identified at the N-terminus of many members belonging to family 19 chitinases, particularly from Classes I and IV (Collinge *et al.*, 1993; Beintema, 1994). The precursor of *Urtica dioica* agglutinin (UDA) consists of two chitin-binding domains at its N-terminus and a C-terminal chitinase domain (Beintema and Peumans, 1992; Lerner and Raikhel, 1992). Following post-translational processing, the UDA precursor is cleaved to generate UDA which lacks the chitinase catalytic domain (Lerner and Raikhel, 1992). Wheat germ agglutinin consists of four chitin-binding domains (Wright *et al.*, 1991). The majority of plant lectins are comprised of hololectins that contain two or more chitin-binding domains which confer agglutination properties (Peumans and Van Damme, 1995).

Since a chitin-binding domain is present in the N-terminus of many members belonging to family 19 chitinases (Collinge *et al.*, 1993), they may be considered as a class of lectins that are structurally linked to an...
unrelated domain (Van Damme et al., 2004). Chitinases are pathogenesis-related proteins that catalyse the random cleavage of internal β-1,4 glycosidic linkages in chitin, a major constituent of fungal cell walls and insect exoskeletons (Boller, 1985). Chitinases exhibit antifungal activity (Schlumbaum et al., 1986) by lysing fungal tips and inhibiting growth (Mauch et al., 1988).

*Brassica juncea* (Indian mustard) BjCHI1 encodes an unusual chitinase with two, almost identical, chitin-binding domains (Zhao and Chye, 1999). Its induced mRNA expression upon wounding, methyl jasmonate treatment, *Aspergillus niger* infection, and caterpillar *Pieris rapae* feeding, suggests it has a role in defence (Zhao and Chye, 1999; Fung et al., 2002). Further, extracts from transgenic tobacco and transgenic potato plants overexpressing BjCHI1 showed activity against *Trichoderma viride* (Chye et al., 2005). Transgenic potato plants were conferred enhanced protection to infection by soil-borne *Rhizoctonia solani* (Fung et al., 2002; Chye et al., 2005). BjCHI1 showed strongest (50–60%) homology to the catalytic domains of family 19 (class I) enzymes, but was structurally distinct from other plant chitinases by the presence of a second chitin-binding domain (Zhao and Chye, 1999). BjCHI1 and UDA are the only known plant proteins that contain two N-terminal chitin-binding domains in tandem. Using *Pichia* expressed recombinant BjCHI1, it was demonstrated that its chitin-binding domains confer agglutination of rabbit erythrocytes, a property first reported in chitinases (Tang et al., 2004). Both chitin-binding domains were deemed essential for erythrocyte agglutination, since this was notably absent in derivatives of BjCHI1 lacking one and both chitin-binding domains. In contrast, the removal of one or both chitin-binding domains in BjCHI1 did not adversely affect chitinase activity (Fung et al., 2002; Tang et al., 2004), consistent with findings from site-directed mutagenesis analysis (Tang et al., 2004) and X-ray crystallography of the catalytic domain (Ubbayasekara et al., 2007).

In this study, BjCHI1 was tested against some common phytopathogens to establish if BjCHI1 could be further applied in crop protection. The phytopathogens tested included *Colletotrichum truncatum*, *C. acutatum*, *Botrytis cinerea*, and Ascochyta rabiei. *Colletotrichum* is the causative agent of anthracnose, a worldwide disease affecting fruit and plantation crops, including papaya, strawberry, tomato, kiwi, pepper, peach, grape, and pecan. *B. cinerea* causes blight (grey mould) and infects many ornamental crops including bulbs and soft fruit crops like grape and strawberry. Ascochyta rabiei causes ascochyta blight of chickpea. The bacterial pathogen *Ralstonia solanacearum* was also tested since it causes bacterial wilt on many tropical plants like banana, tomato, pepper, eggplant, and potato. Finally, the use of BjCHI1 in plastid transformation was explored by testing transplastomic tobacco extracts for chitinase activity followed by bacterial agglutination assays.

**Materials and methods**

**Construction of yeast expression plasmids**

BjCHI1 (Zhao and Chye, 1999; GenBank accession no. AAF02299) and its deletion derivatives lacking one or both chitin-binding domains were cloned into the *Pichia pastoris* expression vector pPIC9K (Invitrogen, San Diego, CA, USA), in-frame to the pPIC9K N-terminal secretory signal peptide (Tang et al., 2004). Polymerase chain reaction (PCR) was used to generate the DNA fragments corresponding to BjCHI1 and its derivatives for expression of recombinant proteins that would be secreted into the growth medium (Tang et al., 2004).

**Production of secreted chitinases expressed in *P. pastoris***

Transformation of *P. pastoris* strain KM71 was carried out by electroporation (Tang et al., 2004). Transformants were verified using PCR and Southern blot analyses before use in protein expression according to the instructions specified in the Multi-copy *Pichia Expression Kit* (Invitrogen). Protein production in *Pichia* was initiated by inoculating an overnight culture grown at 30 °C in 10 ml of BMGY medium [1% (w/v) yeast extract, 2% (w/v) peptone, 100 mM potassium phosphate, pH 6.0, 1.34% (w/v) yeast nitrogen base, 4×10⁻⁵% (w/v) biotin, 1% (w/v) glycerol], to 800 ml of BMGY medium until the optical density (OD) at 600 nm reached at least 2.0. Subsequently, the cells were harvested by centrifugation at 5000 g for 5 min. The cell pellet was then resuspended in 80 ml of BMMY medium [1% (w/v) yeast extract, 2% (w/v) peptone, 100 mM potassium phosphate, pH 6.0, 1.34% (w/v) yeast nitrogen base, 4×10⁻⁵% (w/v) biotin, 0.5% (w/v) methanol] and grown at 30 °C for a further 2 d, during which time methanol was added to a final concentration of 0.5% (v/v) every 24 h. Cells were then harvested by centrifugation at 5000 g for 15 min at 4 °C. The *Pichia*-expressed protein in the supernatant was precipitated overnight using 65% (w/v) ammonium sulphate, following which the precipitated protein was resuspended in 10 mM sodium phosphate buffer, pH 7.2, and dialysed against the same buffer. The dialysed sample was centrifuged at 5000 g at 4 °C for 15 min, filtered through a membrane filter (0.2 μm, Nalgene) and concentrated using a Centricron concentrator (10 kDa MW cut-off; Millipore). The protein was purified on a gel filtration Superdex HiLoad 16/60 column in a Fast Protein Liquid Chromatography (FPLC) system (Amersham Pharmacia Biotechnology, Uppsala, Sweden) according to Tang et al. (2004). Fractions (2 ml), were collected and analysed by sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE; Laemmli, 1970) followed by staining with Coomassie Brilliant Blue to check purity. The purest fractions containing the desired band were pooled and stored at −80 °C.

Protein concentrations were determined using the method of Bradford (1976). These solutions and the control (BSA in 10 mM sodium phosphate buffer, pH 7.2) were filter (0.22 μm) sterilized before use for *in vitro* tests. *Pichia*-expressed crude proteins and FPLC-purified recombinant proteins were loaded on to SDS-PAGE to check protein quality after the determination of protein concentration and before further use for *in vitro* tests.

**Anti-fungal activities using plate growth inhibition assays**

Anti-fungal activities of *Pichia*-expressed, filter-sterilized, FPLC-purified recombinant BjCHI1 and its derivative with one chitin-binding domain were tested *in vitro* using potato dextrose agar (PDA) plates following the method of Fung et al. (2002).
Phytopathogens *C. truncatum* and *C. acutatum* were each inoculated at the centre of the plate and incubated in the dark at 28 °C. Subsequently, various amounts of filter-sterilized recombinant protein (5 µg or 20 µg) dissolved in sodium phosphate buffer (10 mM, pH 7.2) in a total volume of 100 µl, were added to wells 0.5 cm from the growing hyphal front. The plates were placed in the dark at 28 °C and photographed after 72 h. *C. acutatum* and *B. cinerea* were tested using larger amounts of filter-sterilized BjCHI1 (10, 20, 30, 40, or 50 µg). For *C. acutatum*, plates were incubated in the dark at 28 °C followed by photography after 48 h and for *B. cinerea*, plates were incubated in the dark at 20 °C followed by photography after 24 h. *A. rabiei* was inoculated on the centre of a PDA plate for 10–12 d following which *BjCHI1* (0, 4, 8, 12, 16, and 20 µg), dissolved in sodium phosphate buffer (10 mM, pH 7.2) in a total volume of 100 µl, were added to wells 0.5 cm from the growing hyphal front. The plate was placed in the dark at 20 °C and photographed after 5 d. Plate assays were repeated at least five times with reproducible results.

**Determination of IC50 Value**

Antifungal activity of BjCHI1 was expressed as the IC50 value of *C. truncatum*. Hyphal growth inhibition assays (Brockert et al., 1989) were used to determine the concentration required for 50% growth inhibition (IC50). Spore suspensions (500 µl of a stock solution of 2 × 10^6 spores ml^-1) in potato dextrose broth were inoculated in flat bottom wells of a microtitre plate at room temperature for 2 h. BjCHI1 was added to a final concentration of 0, 0.5, 1, 2, 4, 8, or 16 µM ml^-1 and the plates were placed at room temperature until the control germlings (inoculated in broth lacking BjCHI1) attained an average length of 500 µm. The average length of 50 individual hyphae from each well was determined from photomicrographs observed with an inverted microscope. The IC50 value was estimated from the length of the hyphae. Relative hyphal growth was expressed as a percentage of the hyphal growth of control cultures incubated in media lacking BjCHI1. Altogether, three independent experiments were conducted with at least 50 samples taken from each.

**Anti-bacterial assays**

Bacteria tested in anti-bacterial assays were *E. coli*, *R. solanacearum*, *Pseudomonas aeruginosa*, *Micrococcus luteus*, and *Bacillus megaterium*. Inhibition assays on bacterial growth (Larrick et al., 1993) were initiated by inoculation of a single bacterial colony in 50 ml of Luria Broth (for *E. coli*, *P. aeruginosa*, *M. luteus*, and *B. megaterium*) or TTC medium (for *R. solanacearum*). TTC medium contained 0.25% (w/v) dextrose, 1% (w/v) peptone, 0.1% (w/v) Casamino acid (Difco), and 0.005% (w/v) triphenyl tetrazolium chloride. The cultures were grown overnight at 37 °C for *E. coli* and *P. aeruginosa*, 32 °C for *R. solanacearum*, 30 °C for *M. luteus*, and 28 °C for *B. megaterium*. Each overnight culture (50 µl) was further diluted 1:1000 using 50 ml fresh medium to a final concentration of 10^5 colony-forming units ml^-1. A microtitre well containing 80 µl of diluted bacterial culture (10^5 colony-forming units ml^-1), BjCHI1, its deletion derivatives or BSA (control) was added to obtain final concentrations of 10–60 µg ml^-1. Sodium phosphate buffer (10 mM, pH 7.2) was added to a total volume of 100 µl. An OD600 reading was recorded after 30 min and a second reading was taken following incubation for 24 h at the appropriate incubation temperatures. An increase in OD600 would indicate bacterial growth. Three independent experiments, with three replicates in each, were performed altogether.

**Bacterial agglutination assays**

Agglutination assays were tested using Gram-negative bacteria (*E. coli*, *R. solanacearum*, *P. aeruginosa*) and Gram-positive bacteria (*M. luteus* and *B. megaterium*). In bacterial agglutination assays (Saito et al., 1995), 25 µl suspension of each bacterial culture (OD600 = 10) was added to 25 µl of BjCHI1 solution to obtain final concentrations of 0, 7.8, 15.6, 31.3, 62.5, 125, and 250 µg ml^-1, in wells on a microtitre plate, and left for 24 h at room temperature. Agglutinating activities were observed and expressed as the minimum agglutinating concentration (Saito et al., 1995).

To investigate the inhibition of BjCHI1 binding activity by free monosaccharides (galactose and glucose) and disaccharides (lactose), 25 µl of BjCHI1 solution (at a final concentration of 250 µg ml^-1) was preincubated with 50 µl of galactose, glucose or lactose, at 37 °C for 1 h, before addition of 25 µl of a *R. solanacearum* culture. The mixtures, in wells of a microtitre plate, were incubated at room temperature for 24 h. Each sugar was tested at final concentrations of 0, 125, 625, 12.5, 25, 50, and 100 mM. The inhibition of BjCHI1 binding activity by each monosaccharide or disaccharide was expressed as the minimum inhibitory concentration (Hirata et al., 1990).

The agglutinating activity of purified chitinase from transplastomic tobacco lines was similarly tested against *R. solanacearum*. Affinity column-purified BjCHI1 chitinase from transplastomic tobacco was used at final concentrations of 0, 125, 250, and 500 µg ml^-1 in wells on a microtitre plate, and left to incubate with *R. solanacearum* at room temperature for 24 h.

**Construction of plastid transformation vector for BjCHI1 expression**

Plastid transformation vector pMLVHisBj was constructed from plasmid pMLVHisA (Zhou et al., 2006), a derivative of pVSR326 (Reddy et al., 2002). Plasmid pVSR326 utilizes the tobacco plastid genome sequences spanning *rbcL–accD* to target the reporter gene encoding β-glucuronidase (*GUS*) into the chloroplast genome by homologous recombination (Reddy et al., 2002). The promoter and terminator for *GUS* are derived from the rice plastid gene *psbA*, which encodes a 32 kDa photosystem II protein. The selectable marker gene *aadA*, which specifies spectinomycin-resistance, is expressed from the promoter of the rice plastid 16S rRNA operon (*rrn*) and *aadA* is located adjacent to *GUS* (Reddy et al., 2002). In the pMLVHisA vector, transcription of the transgene is under the control of the rice plastid *psbA* promoter and the *psbA* terminator. In the construction of pMLVHisA, other than unique restriction sites introduced to pSRR326 to facilitate the cloning of the transgene for transplastomic expression, a start codon plus a (His)6-tag was incorporated to tag the recombinant protein. This enables recognition of the recombinant protein using antiserum against the (His)6-tag in western blot analysis and easy purification of the recombinant protein, if and when required, using Ni-NTA Agarose (Qiagen) affinity columns.

A 1.3 kb *ApaI* fragment encoding BjCHI1, including 7 bp of the 5′-untranslated region, was cloned into the *ApaI* site of pMLVHisA. In the resultant derivative, designated pMLVHisBj, amino acids 1-393 of BjCHI1 were fused in-frame to 18 amino acids derived residues, including the ‘ATG’ start codon and the (His)6-tag. The presence of flanking *rbcL* and *accD* sequences from the tobacco plastid genome enabled the incorporation of BjCHI1 (and *aadA*) into the tobacco plastid genome via homologous recombination.

**Tobacco plastid transformation**

Sterilized tobacco seeds were germinated on Murashige and Skoog (MS) medium (Murashige and Skoog, 1962), supplemented with agar (5 g l^-1) and sucrose (30 g l^-1) for about 6 weeks to obtain the leaf material for transformation. Plastid transformation (Swab et al., 1993) was carried out by bombardment of tobacco leaves with tungsten particles (M 17) coated with pMLVHisBj plasmid DNA.
using the particle delivery system, PDS 1000-He (Bio-Rad) and its
accessories. For bombardment, leaves were placed abaxial side up
on RMOP medium consisting of MS salts, 6-benzylaminopurine
(1 mg l⁻¹), α-naphthalene acetic acid (0.1 mg l⁻¹), thiamine (1 mg
l⁻¹), myo-inositol (100 mg l⁻¹), agar (5 g l⁻¹) at pH 5.7, and
sucrose (30 g l⁻¹). Following bombardment, leaves were cut into
small pieces and placed on RMOP selection medium containing
spectinomycin dihydrochloride (500 mg l⁻¹). The regenerated
tobacco plantlets obtained following plastid transformation were
subjected to five more cycles on selection medium to obtain
homoplastomic plastid-containing plants.

Plantlets were tested by PCR analysis using a primer pair
consisting of a BjCHI1-specific primer and a pMLVHisBj vector-
specific primer to confirm integration of an intact insert via
homologous recombination. Subsequently, a pair of primers that
flank the inserted DNA in the tobacco plastome was used in PCR
to estimate the percentage of homoplastomic genomes in each trans-
plastomic lines. Each PCR reaction (25 µl) consisted of 250 ng
template, 5 pmol of each primer, 0.75 U AmpliTaq Gold (Applied
Biosystems), 2.5 µl 10× PCR reaction buffer, 1.5 µl of 25 mM
MgCl₂, and 0.25 µl of each 10 mM dNTP. PCR-amplification was
initiated with denaturation at 95 °C for 3 min, followed by 35
cycles of 95 °C for 1 min, 58 °C for 1 min, and 72 °C for 1 min.

Southern blot, northern blot, and western blot analyses on
transplastomic tobacco lines
Following preliminary analysis by PCR, the transplastomic tobacco
lines were tested using Southern blot, northern blot, and western
blot analyses. For Southern blot analysis, 20 µg DNA isolated
according to Dellaporta et al. (1983) was digested with BamHI,
separated by electrophoresis in 0.7% agarose gel, and blotted onto
Hybond-N (Amersham) membranes (Sambrook et al., 1996).
Membranes were prehybridized in a solution containing
50% Hybond-N (Amersham) membranes (Sambrook et al., 1996),
following preliminary analysis by PCR, the transplastomic tobacco
plantlets obtained following plastid transformation were
subjected to five more cycles on selection medium to obtain
homoplastomic plastid-containing plants.

Sambrook et al. according to Dellaporta et al. (1983) was digested with BamHI,
separated by electrophoresis in 0.7% agarose gel, and blotted onto
Hybond-N (Amersham) membranes (Sambrook et al., 1989).
Membranes were prehybridized in a solution containing
50% denatured formamide, 6× SSC, 5× Denhardt’s, 1% SDS, 100 mg
ml⁻¹ denatured salmon sperm DNA at 42 °C for 4–6 h. Random-
primed 3²P-labelled rbcL probe was added and hybridized over-
night. Membranes were washed in 0.1× SSC, 0.1% SDS at 65 °C.

For northern blot analysis, RNA (20 µg) extracted from whole
plants (Nagy et al., 1988), was denatured at 50 °C for 30 min in
glyoxal, separated by electrophoresis in 1.5% agarose gel, and
transferred to Hybond-N membrane (Amersham). Blots, prehybrid-
dized for 4–6 h, were hybridized with random-primed 3²P-labelled
BjCHI1 cDNA (Zhao and Chye, 1999) in a solution containing 50%
denatured formamide, 1× Denhardt’s, 6× SSPE (1× SSPE: 0.15 M NaCl,
0.01 M NaH₂PO₄, 0.001 M EDTA, pH 7.4), 0.1% SDS, 100 µg
ml⁻¹ denatured salmon sperm DNA, and 10% dextran sulphate at
42 °C overnight. Blots were washed in 0.1× standard saline citrate
(1× SSC:150 mM NaCl, 15 mM sodium citrate, pH 7.0) containing
0.1% SDS at 65 °C.

For western blot analysis, total plant protein was prepared from
whole plants according to Kush et al. (1990). Twenty µg of total
protein was separated by SDS-PAGE and transferred onto Hybond-C
(Amersham) membrane as described by Sambrook et al. (1989). In
western blot analysis, cross-reacting bands were detected using
polyclonal antibodies against BjCHI1 following the procedures
described in the Amplified Alkaline Phosphatase Goat Anti-Rabbit
Immuno-Blot Assay Kit (Bio-Rad). Polyclonal antibodies were raised
in rabbits against a synthetic peptide (YKEEIDKSDPHC) corre-
sponding to amino acids 231–242 of BjCHI1 (Zhao and Chye, 1999).

Chitinase assays
For colorimetric chitinase assays (Wirth and Wolf, 1990) using the
substrate carboxymethylchitin-Remazol Brilliant Violet 5R (CM-
chitin-RBV; Loewe Biochemica GmbH, Sauerlack, Germany),
samples of plant extracts (1–10 µg) were made up to 250 µl with
water and incubated with 250 µl 0.1 M sodium acetate buffer, pH
5.0 and 250 µl CM-RBV-chitin at 37 °C for 1 h. The reaction was
stopped with 250 µl 1 M HCl, placed on ice for 15 min to
precipitate the non-degraded chitin, and the reaction mixture
subsequently centrifuged at 20 000 g at 4 °C for 5 min. OD₅₅₀nm
was read and values were corrected for background, which was
determined by using values obtained with protein that had been
heated for 10 min at 100 °C to destroy activity. Three replicates
were performed and average values used. One unit of chitinase was
defined as the enzyme activity liberating 1 mg of soluble N-
acyetylglucosamine (GlnNAc) equivalent h⁻¹ at infinite dilution.

Results

BjCHI1 inhibits growth of C. truncatum, C. acutatum,
B. cinerea, and A. rabiei in vitro
Antifungal activities of BjCHI1 and its deletion derivative
with one chitin-binding domain were demonstrated using
radial growth-inhibition assays (Schlumbaum et al., 1986). Growth inhibition of C. truncatum (Fig. 1A) and
C. acutatum (Fig. 1B) were observed in wells to which
BjCHI1 (wells 2 and 3, containing 5 µg and 20 µg,
respectively) and its derivative (wells 4 and 5, containing
5 µg and 20 µg, respectively) were added, in contrast to
wells containing buffer control (well 1) in which fungal
hyphae had grown within the wells.

When the amounts of BjCHI1 were increased beyond 20
µg, significant inhibition of C. acutatum was observed in
wells containing 30 µg (Fig. 1C, well 4), 40 µg (Fig. 1C,
well 5), and 50 µg (Fig. 1C, well 6) of BjCHI1. Similar
results were evident in plate assays on B. cinerea (Fig. 1D)
at which BjCHI1 at 0 µg (well 1), 10 µg (well 2), 20 µg
(well 3), 30 µg (well 4), 40 µg (well 5), and 50 µg (well 6)
were tested. Figure 1E shows results from plate assays on
A. rabiei using BjCHI1 at 0 µg (well 1), 4 µg (well 2), 8 µg
(well 3), 12 µg (well 4), 16 µg (well 5), and 20 µg (well 6).
A. rabiei was noticeably inhibited by BjCHI1 at 20 µg
(Fig. 1E, well 6).

These results demonstrate the in vitro activity of
BjCHI1 against phytopathogens C. truncatum, C. acutatum,
B. cinerea, and A. rabiei. Results of anti-fungal activities on C. truncatum and C. acutatum exhibited by
BjCHI1 and its derivative with one chitin-binding domain are
consistent with observations of these two proteins on
Trichoderma viride in plate assays (Fung et al., 2002).
BjCHI1 and its derivatives lacking one or both chitin-
binding domains were also observed to inhibit spore
germination of T. viride (data not shown), confirming
previous observations that chitinase activity is unaffected
by removal of either one or both chitin-binding domains
(Fung et al., 2002; Tang et al., 2004).

BjCHI1 inhibits spore germination of C. truncatum in
vitro
The hyphal growth inhibition assay was used to
determine the IC₅₀ value of BjCHI1 against phytopathogen
Growth of hyphae in BjCHI1-containing medium was examined in three independent experiments. The $IC_{50}$ value of BjCHI1 on *C. truncatum* was estimated to be approximately 1 µg ml$^{-1}$. Germination of spores grown in potato dextrose broth as observed by light microscopy, was minimal (~15% relative hyphal growth) after 24 h of incubation in potato dextrose broth in the presence of BjCHI1 at 8 µg ml$^{-1}$. By contrast, the average hyphal length of the spores incubated in a broth lacking BjCHI1 reached an average of 500 µm.

**Pichia-expressed recombinant BjCHI1 shows antibacterial activity in vitro**

When BjCHI1 and its derivatives lacking one or both chitin-binding domains were tested against *E. coli*, *R. solanacearum*, *P. aeruginosa*, *M. luteus*, and *B. megaterium*, only BjCHI1 was observed to inhibit the growth of all bacteria tested. In these tests, BSA was used as a control. Distinct BjCHI1-mediated inhibition ($P < 0.1$) was observed at various concentrations with different strains, for example, 20 µg ml$^{-1}$ for *E. coli* and *R. solanacearum*, but at higher BjCHI1 concentrations for other bacteria (Fig. 2). In contrast, both BjCHI1 derivatives did not show significant inhibition using the Student’s $t$ test.

A comparison of growth inhibition using BjCHI1 at 60 µg ml$^{-1}$ for all bacteria tested revealed that *E. coli* was most susceptible (Fig. 2), followed by *R. solanacearum* and *P. aeruginosa*. The inhibition of *M. luteus* was less significant, and it was negligible for *B. megaterium*. The most BjCHI1-susceptible bacteria (*E. coli*, *R. solanacearum*, and *P. aeruginosa*) are Gram-negative, while the other least BjCHI1-susceptible bacteria (*M. luteus* and *B. megaterium*) are Gram-positive.

**BjCHI1 agglutinates Gram-negative bacteria**

Since the data from bacterial growth inhibition assays suggested that BjCHI1 was more effective on Gram-negative bacteria, which differed in cell wall composition from Gram-positive bacteria, further investigations were then carried out to check if inhibition was related to BjCHI1-mediated bacterial agglutination.

---

**Fig. 1.** BjCHI1 inhibits growth of phytopathogens *Colletotrichum truncatum*, *C. acutatum*, *Botrytis cinerea*, and *Ascochyta rabiei*. (A, B) Photographs of the bottom of Petri plates showing the effect of BjCHI1 and its derivative with one chitin-binding domain on the growth of *C. truncatum* (A) and *C. acutatum* (B). FPLC-purified proteins were added in wells when the hyphal front was 0.5 cm from the wells. Well 1, control with 10 mM sodium phosphate buffer (pH 7.2); well 2, 5 µg BjCHI1; well 3, 20 µg BjCHI1; well 4, 5 µg BjCHI1 derivative, and well 5, 20 µg BjCHI1 derivative. The photograph was taken 72 h after addition of protein. (C, D) Photographs of the top of the plates showing the effect of BjCHI1 on the growth of *C. acutatum* (C) and *B. cinerea* (D). Well 1, control with 10 mM sodium phosphate buffer (pH 7.2); well 2, 10 µg BjCHI1; well 3, 20 µg BjCHI1; well 4, 30 µg BjCHI1; well 5, 40 µg BjCHI1 and well 6, 50 µg BjCHI1. (E) Photograph of the top of the plate showing the effect of BjCHI1 on the growth of *A. rabiei*. Well 1, control with 10 mM sodium phosphate buffer (pH 7.2); well 2, 4 µg BjCHI1; well 3, 8 µg BjCHI1; well 4, 12 µg BjCHI1; well 5, 16 µg BjCHI1, and well 6, 20 µg BjCHI1.

**Fig. 2.** BjCHI1-mediated growth inhibition of bacteria. Inhibition curves of BjCHI1 on *B. megaterium*, *M. luteus*, *P. aeruginosa*, *R. solanacearum*, and *E. coli*. Relative bacterial growth (%) in the presence of varying amounts of BjCHI1 in comparison to BSA (100%). The results were derived from three independent experiments (each carried out in triplicate) and are mean values + SD.
Agglutination assays were carried out and agglutinating activities were observed and expressed as the minimum agglutinating concentration (MAC). These assays on *E. coli*, *R. solanacearum*, and *P. aeruginosa* revealed that higher concentrations of BjCHI1 culminated in greater agglutination (Fig. 3A). By contrast, agglutination was absent for *M. luteus* and *B. megaterium* with increasing amounts of BjCHI1 (Fig. 3A). The MAC values (Table 1) when compared to the BSA control were consistent with results from bacterial growth inhibition. Enhanced agglutination of *E. coli* and *R. solanacearum* was correlated with stronger inhibition (56% and 40%, respectively, in Table 1) as shown in the bacterial growth inhibition assays in Fig. 2. The slight self-agglutination of *P. aeruginosa* resulting from basal agglutination could account for the comparably lower degree of inhibition seen in the bacterial growth inhibition assays. Lack of agglutination with Gram-positive bacteria *M. luteus* and *B. megaterium* was consistent with results of a lack of BjCHI1-mediated growth inhibition for *M. luteus* and *B. megaterium* (Table 1).

**Free monosaccharides or disaccharides relieve binding activity of BjCHI1**

Inhibition of carbohydrate-binding activity was performed following Hirata *et al.* (1990). Three types of carbohydrates (galactose, glucose, and lactose) were tested, and each result was expressed as the minimum inhibitory concentration against 125 μg ml\(^{-1}\) of BjCHI1. Since *R. solanacearum* demonstrated the most consistent agglutination response on addition of BjCHI1, it was used as the sample bacterium in the agglutination inhibition assays. Progressive inhibition of agglutination was observed when the added amount of carbohydrate was increased (Fig. 3B–D). Both free galactose and glucose inhibited agglutination of *R. solanacearum*, although the latter displayed a slightly stronger inhibitory effect (Fig. 3B, C). The result is consistent with the structure of lipopolysaccharide, of which the outer core contains a pentasaccharide of glucose, galactose, and *N*-acetylglucosamine, and the *O*-antigen consists of mannose (with sugar specificity similar to glucose), and rhamnose, amongst others. Remarkably, most chitin-binding lectins containing hevein domains are *N*-acetylglucosamine-specific (Beintema, 1994). This provides an insight on the mechanism through which BjCHI1 agglutinates Gram-negative bacterial cells.

**Analyses on transplastomic tobacco lines expressing BjCHI1**

Following preliminary PCR analysis using BjCHI1-specific and plastid vector-specific primer pairs, as well as primer pairs flanking the inserted DNA in the tobacco plastome, Southern blot analysis was used to confirm the integration of BjCHI1 by digestion of DNA from the transplastomic tobacco lines, with *Bam*HI. Southern blot analysis was carried out with wild-type tobacco as a control by using a 32P-labelled rbcL probe. The results (Fig. 4A) showed the presence of a 1.2 kb hybridizing band in wild-type (lane 1) tobacco and a strong 3.4 kb hybridizing band in the two transplastomic tobacco lines (His-BjCHI1-3 in lane 2 and HisBjCHI1-4 in lane 3) confirming integration of the BjCHI1 cDNA in the tobacco plastome of these transplastomic lines. The 3.4 kb band was clearly absent in wild type (Fig. 4A, lane 1). The 1.2 kb hybridizing rbcL BamHI band corresponds to the BamHI fragment (nucleotide positions 58051–59288) within rbcL in the tobacco plastome (Accession No. Z00044). The 3.4 kb BamHI recombinant band resulting from homologous recombination, is due to cleavage corresponding to the BamHI restriction sites at rbcL.

![Fig. 3. BjCHI1-mediated agglutination of bacteria. (A) Bacteria, *E. coli*, *R. solanacearum*, *P. aeruginosa*, *M. luteus*, and *B. megaterium*, were tested using BjCHI1 at final concentrations of 0 (well 1), 7.8 (well 2), 15.6 (well 3), 31.3 (well 4), 62.5 (well 5), 125 (well 6), and 250 (well 7) μg ml\(^{-1}\). (B–D) Effect of galactose (B), glucose (C) or lactose (D) on the agglutination of *R. solanacearum*. BjCHI1 (125 μg ml\(^{-1}\)) was preincubated with final sugar concentrations at 0 nM (well 1), 3.125 nM (well 2), 6.25 nM (well 3), 12.5 nM (well 4), 25 nM (well 5), 50 nM (well 6), and 100 nM (well 7) before the addition of *R. solanacearum.*](http://jxb.oxfordjournals.org/)

**Table 1. BjCHI1-mediated agglutination and anti-bacterial activity**

<table>
<thead>
<tr>
<th></th>
<th>MAC(^a) (μg ml(^{-1}))</th>
<th>MPC(^b)</th>
<th>RIA(^c) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>E. coli</em></td>
<td>7.8</td>
<td>Self-precipitated</td>
<td>56</td>
</tr>
<tr>
<td><em>R. solanacearum</em></td>
<td>7.8</td>
<td>Self-precipitated</td>
<td>56</td>
</tr>
<tr>
<td><em>P. aeruginosa</em></td>
<td>15.6</td>
<td>Self-precipitated</td>
<td>40</td>
</tr>
<tr>
<td><em>M. luteus</em></td>
<td>&gt; 250</td>
<td>31.25 μg ml(^{-1})</td>
<td>14</td>
</tr>
<tr>
<td><em>B. megaterium</em></td>
<td>&gt; 250</td>
<td>Self-precipitated</td>
<td>4</td>
</tr>
</tbody>
</table>

\(^a\) Minimum agglutinating concentration.
\(^b\) Minimum precipitating concentration.
\(^c\) Relative inhibitory activity (RIA) at 60 μg ml\(^{-1}\); RIA = [1 – the growth of bacteria in 60 μg ml\(^{-1}\) BjCHI1/growth of bacteria in control group] × 100%.
(nucleotide position 58051 in the tobacco plastome) and at \( accD \) (nucleotide position 60656), but inclusive of the pMLVHisBj vector DNA between them. Upon homologous recombination at the plastid flanking sequences \( rbcL \) and \( accD \) regions, these two \( BamHI \) sites are expected to be 3.4 kb apart. In the \( BjCHI1 \) plastid transformation vector pMLVHisBj, the \( BjCHI1 \) cDNA was cloned between the plastid flanking sequences (\( rbcL \) and \( accD \)) to enable \( BjCHI1 \) to be inserted into the tobacco plastome.

Following the confirmation of gene integration into the plastid genome, northern blot analysis was carried out on transplastomic tobacco plants. Results of northern blot analysis (Fig. 4B) using \( 32P \)-labelled \( BjCHI1 \) cDNA showed the presence of a 1.3 kb hybridizing band in both lines (HisBjCHI1-3 in lane 2 and HisBjCHI1-4 in lane 3), whereas it was absent in the wild type (lane 1), confirming efficient transcription of \( BjCHI1 \) from the plastid genome.

Ethidium bromide-stained rRNA bands (Fig. 4B, bottom) showed even loading of total RNA.

Following Southern blot and northern blot analyses, transplastomic tobacco plants were further tested by western blot analysis using anti-\( BjCHI1 \) specific antibodies. Western blot analysis using these antibodies (Fig. 4C) on crude protein from \( BjCHI1 \) transplastomic tobacco (HisBjCHI1-3 in lane 2 and HisBjCHI1-4 in lane 3) showed a cross-reacting \( BjCHI1 \) band with an apparent molecular mass of 37 kDa in the transplastomic lines, while no such band was evident in the wild type (lane 1).

Figure 5A indicates that transplastomic tobacco line HisBjCHI1-4 (0.028 U g\(^{-1}\) min\(^{-1}\)) showed greater than a 2-fold increase in chitinase activity compared to the wild-type tobacco line (0.013 U g\(^{-1}\) min\(^{-1}\)). When agglutinating activity of affinity-purified chitinase from transplastomic tobacco lines against \( R.\ solanacearum \) was examined, a gradual increase of agglutinating activity was observed with an increase in the chitinase final concentration from 0 \( \mu g \) to 500 \( \mu g \) ml\(^{-1}\); clear agglutination activity was observed at the highest concentration tested (Fig. 5B).

**Discussion**

Initial attempts to express (His)\(_6\)-\( BjCHI1 \) in \( E.\ coli \) resulted in problems with poor bacterial growth (Zhao and Chye, 1999). Subsequently, \( BjCHI1 \) and its deletion derivatives lacking one or both chitin-binding domains...
were expressed in *Pichia* for efficient mass production of the protein (Tang et al., 2004). This investigation further shows that *Pichia*-expressed BjCHI1 inhibits growth of Gram-negative bacteria, providing an eventual explanation of its adverse effect on *E. coli* growth in a previous report (Zhao and Chye, 1999). The importance of the second chitin-binding domain in conferring anti-bacterial activity on Gram-negative bacteria is also demonstrated because BjCHI1 derivatives lacking one or both chitin-binding domains did not confer anti-bacterial properties. It has been previously shown that only BjCHI1 with both chitin binding domains (but not its derivatives) has the ability to agglutinate rabbit erythrocytes. This study further showed that BjCHI1 agglutinates the Gram-negative bacterial pathogen, *R. solanacearum*, thus extending anti-fungal action. Since agglutination is largely an extracellular response, the different biochemical nature of the cell wall of Gram-positive and Gram-negative bacteria, could to some extent dictate the interaction involving the duplicated chitin (carbohydrate)-binding domain of BjCHI1. In contrast, the removal of one or both chitin-binding domains did not adversely affect its catalytic activity in chitinase assays using *Pichia*-derived BjCHI1 (Tang et al., 2004). These results are consistent with the agglutination function of other known carbohydrate-binding proteins (Peumans and Van Damme, 1995).

Plant lectins are defined as plant proteins possessing at least one non-catalytic domain, which binds reversibly to a specific mono- or oligosaccharide (Peumans and Van Damme, 1995). Monovalent lectins, possessing only one carbohydrate-binding domain, do not precipitate glyco-conjugates or agglutinate cells. Previous studies have shown that hololectins may effectively agglutinate cells because they contain two or more carbohydrate binding domains which are identical or very homologous (Van Damme et al., 1998). Interestingly, the amino acid sequence of the two chitin-binding domains of BjCHI1 shows 95% identity with each other (Zhao and Chye, 1999). The bivalent structure of BjCHI1 may be a prototype for plant pathogenesis-related proteins that are able to agglutinate and inhibit bacteria cell growth. The precursor of UDA resembles BjCHI1 by the presence of two chitin-binding domains (Zhao and Chye, 1999) and has been classified as a hololectin (Van Damme et al., 1998). However, UDA differs from BjCHI1 because its catalytic domain is eliminated from the two chitin-binding domains during post-translational cleavage. Observation reported herein of inhibition from agglutination by free monosaccharides (galactose and glucose) and disaccharide (lactose) further suggests lectin-lipopolysaccharide interaction between BjCHI1 and Gram-negative bacterial cells.

The IC$_{50}$ value of BjCHI1 on the growth of *C. truncatum* was approximately 1 µg ml$^{-1}$. In comparison, this is lower than most other plant chitinases with an average value of 2 µg ml$^{-1}$. The IC$_{50}$ value against *T. viride* for UDA, which lacks the catalytic domain, is 50 µg ml$^{-1}$ (Broekaert et al., 1989). However, addition of low concentrations of UDA to other tobacco chitinases increased the inhibitory potency of the chitinase by about 4-fold, indicating a synergistic action of UDA and chitinases. UDA enhances accessibility to the active sites with its two chitin-binding domains, and the chitinase confers the catalytic function (Broekaert et al., 1989). BjCHI1, a functional protein that includes both the double chitin-binding domains and the catalytic domain in one molecule, potentially resembles the combinatorial action of UDA acting in synergism with other chitinases. BjCHI1 exhibited combined anti-microbial activity in combating fungi and in conferring anti-bacterial properties in defence against Gram-negative phytopathogens. In the current study, the BjCHI1 derivative with one chitin-binding domain, also showed the ability to inhibit fungal hyphae growth in *in vitro* plate assays against *C. truncatum* and *C. acutatum*, in accordance with previous studies that demonstrated similar levels of chitinase activities in BjCHI1 and this derivative using standard colorimetric chitinase assays (Fung et al., 2002; Tang et al., 2004).

BjCHI1 as a chimera comprising an N-terminal duplicated chitin-binding domains conferring agglutination-mediated anti-bacterial properties and a C-terminal chitinase domain providing anti-fungal activity, is structurally an effective anti-microbial agent. An intriguing question relates to the presence of the second chitin-binding domain in BjCHI1, which may be redundant in antifungal activity, as the majority of chitinases consist of at most only one chitin-binding domain and a catalytic domain (Collinge et al., 1993). Based on the non-antibacterial nature of the BjCHI1 derivatives lacking chitin-binding domains, the bivalent structure of the carbohydrate-binding domains in BjCHI1 now appears especially significant in defence against bacteria. Molecular and structural studies have shown that cross-linking interactions between certain bivalent and multivalent lectins result in the formation of unique two- and three-dimensional supermolecular assemblies (Brewer, 1997). Mobile infecting bacterial cells could be immobilized and inhibited by such supermolecular assemblies as a result of BjCHI1 binding bacterial cells. In addition, the appearance of a second chitin-binding domain may enhance the ability of BjCHI1 to access the reactive sites on fungal hyphae.

Chitinolytic activity against insects, nematodes, and fungi raises the use of chitinases as alternative pesticides which are environmentally-friendly compared to chemical pesticides. In this study, BjCHI1 showed dual roles as a defence protein against both pathogenic bacteria, including *R. solanacearum*, as well as fungal phytopathogens, *C. truncatum*, *C. acutatum*, *B. cinerea*, and *A. rabiei*. Given that extracts from BjCHI1-overexpressing transplastomic tobacco conferred both chitinase and agglutination activities, BjCHI1 could be potentially promising for future applications in plant protection in agriculture. However,
despite the many advantages that plastid transformation has over nuclear transformation in the expression of foreign proteins, this relatively new technology is applicable to only a few plant species (Maliga, 2004). Perhaps, ultimately a combination of nuclear and plastid transformation approaches using multiple genes may be necessary to attain disease resistance in genetically-engineered plants. Limitations encountered in the production of disease-resistant plants also arise from the diversity in phytopathogens, ranging from bacteria, and fungi to algal Oomycetes, and the complexities in their lifestyles, which may involve biotrophy, hemibiotrophy, and necrotrophy, making it difficult for any one approach to be effective (Collinge et al., 2008). Indeed, useful levels of protection have not been accomplished from strategies involving single pathogen- or related proteins (Gurr and Rushton, 2005; Collinge et al., 2008). Perhaps the use of synthetic ‘designer’ and ‘lifestyle-specific’ promoters may be necessary to alleviate some of these problems currently hindering genetic engineering in disease protection (Gurr and Rushton, 2005).

Acknowledgements

We thank Professor CY Ma for the provision of FPLC facilities. This work was supported the University of Hong Kong (10205761 and 10208034). MLC was recipient of a Dyason Universitas 21 Fellowship at the University of Melbourne.

References


