Bacterial chitinase with phytopathogen control capacity from suppressive soil revealed by functional metagenomics

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Abstract Plant disease caused by fungal pathogens results in vast crop damage globally. Microbial communities of soil that is suppressive to fungal crop disease provide a source for the identification of novel enzymes functioning as bioshields against plant pathogens. In this study, we targeted chitin-degrading enzymes of the uncultured bacterial community through a functional metagenomics approach, using a fosmid library of a suppressive soil metagenome. We identified a novel bacterial chitinase, Chi18H8, with antifungal activity against several important crop pathogens. Sequence analyses show that the \( \text{chi18H8} \) gene encodes a 425-amino acid protein of 46 kDa with an N-terminal signal peptide, a catalytic domain with the conserved active site F\(^{175}\)DGIDIDWE\(^{183}\), and a chitinase insertion domain. Chi18H8 was expressed (pGEX-6P-3 vector) in \( \text{Escherichia coli} \) and purified. Enzyme characterization shows that Chi18H8 has a prevalent chitobiosidase activity with a maximum activity at 35 °C at pH lower than 6, suggesting a role as exochitinase on native chitin. To our knowledge, Chi18H8 is the first chitinase isolated from a metagenome library obtained in pure form and which has the potential to be used as a candidate agent for controlling fungal crop diseases. Furthermore, Chi18H8 may also answer to the demand for novel chitin-degrading enzymes for a broad range of other industrial processes and medical purposes.

Keywords Soil metagenome · Metagenomic library · Chitinolytic enzyme · Phytopathogen control · Antifungal · Heterologous protein expression

Introduction

Disease of plants caused by fungal pathogens contributes to extensive loss of crops important for food and energy production globally. Moreover, the norm of monoculture practice, further increases opportunities for the invasion of phytopathogens (Herrera-Estrella and Chet 1999). As a consequence, the use of synthetic fungicides, many which are toxic, is extensive and evidently results in costs for public and ecosystem health (Mullin et al. 2010). A more environmentally sustainable approach to the use of toxic chemicals is microbiological control of fungal disease employing bacteria that exhibit antifungal action (Herrera-Estrella and Chet 1999). There are soils that are naturally suppressive toward plant diseases and microorganisms in these soils are proposed to be involved in the suppressiveness (Borneman and Becker 2007; Steinberg et al. 2007). As a result, several bacterial species have been isolated and commercially introduced for biocontrol purposes (Steinberg et al. 2007 and references therein). However, given the inherent limitations in the use of living organisms, such as relatively short shelf life of the products or inconsistent performance in the field,
their application is confined (Neeraja et al. 2010). Alternative solutions are formulations with enzymes possessing antiphytopathogenic activity.

Promising candidates for this purpose are bacterial chitinases, as these degrade chitin, one of the main constituents of the fungal cell wall (Herrera-Estrella and Chet 1999; Zhang et al. 2001). Chitinolytic enzymes are also generally attractive for their potential use in a broad range of biotechnological applications, for example in biofuel production or bioconversion processes on shellfish waste to obtain value-added products, such as chitosan and chitooligosaccharides for the pharmaceutical market (Bhattacharya et al. 2007; Li and Greene 2010). In an ecological perspective, bacterial chitinases are crucial in the global biogeochemical re-cycling of carbon and nitrogen through the hydrolyzation of chitin. After cellulose, chitin is the most abundant biopolymer in nature, widely distributed within exoskeletons invertebrates, fungal cell walls, marine diatoms, crustaceans, and zooplankton (Gooday 1990). It is otherwise rather resistant to degradation and would without bacterial chitinases be trapped in biomass as insoluble in nature (for reviews, see Karlsson and Stenlid 2009; Keyhani and Roseman 1999). Degradation of chitin enables bacterial utilization of the end products, chitobiose, and $N$-acyethylglucosamine compounds, as an energy-, carbon-, and/or nitrogen source (Gooday 1990).

Most of the bacterial chitinases are glycosyl hydrolases of family 18 (Henrissat and Bairoch 1993), which can be further classified into subfamilies A to C based mainly on amino acid sequence similarities of the catalytic domain and a conserved consensus motif of the catalytic site (Henrissat and Bairoch 1993; Karlsson and Stenlid 2009; Suzuki et al. 2002). Depending on the catalytic specificity as a result of enzyme structure, chitinases may show either endo- or exo-activity (Henrissat and Davies 2000). To date, bacterial chitinase genes have been identified, by conventional molecular screening approaches, in bacterial isolates or uncultured bacteria within both aquatic and soil environments (for example, Hobel et al. 2005; Ikeda et al. 2007; LeCleir et al. 2004; Metcalf et al. 2002; Ramaiah et al. 2000; Terahara et al. 2009; Uchiyama and Watanabe 2006). Only a few studies, however, have used metagenomics tools to identify novel bacterial chitinase genes (Cottrell et al. 1999; LeCleir et al. 2007). The advantage of metagenome-based approaches is the complete access to the entire community genetic makeup without the need of microbial cultivation (for review, see Sjöling et al. 2007). To our knowledge, none of these studies has yet resulted in the isolation and characterization of novel biologically active chitinases.

As chitin degradation is such an important environmental function, we investigated, in a previous study, the effect of chitin amendment to a suppressive field soil on the bacterial community structure (Hjort et al. 2007). We could show using terminal restriction fragment length polymorphism and denaturing gradient gel electrophoresis analyses—that chitin amendment to the soil dramatically increased the relative abundances of known chitin-degrading genera, such as Oerskovia, Kitaatospora, and Streptomyces. These organisms became dominant also among the actively growing bacteria in the community (Hjort et al. 2007). Given that the soil bacterial community of the suppressive field responded to chitin amendment (Hjort et al. 2007), the active community contained taxa that typically are chitinolytic (Hjort et al. 2007) and that a number of isolates with antifungal and chitinase activity previously were obtained from this soil (Adesina et al. 2007), we sought to investigate this soil metagenome for novel chitinolytic enzymes with biocontrol capacity, suitable for more environmentally sustainable applications in agricultural processes. In this study, we used a targeted screening approach based on genetic and functional metagenomics with the aim to identify novel chitinolytic enzymes through heterologous expression. Our aim was to characterize further isolated enzymes for the purpose of identifying a promising candidate for phytopathogen biocontrol.

Materials and methods

Metagenomic library

A soil characterized as suppressive to club-root disease of cabbage (Worku and Gerhardson 1996) was sampled from a field at the Swedish University of Agricultural Sciences in Uppsala, Sweden (Hjort et al. 2007), and a fosmid metagenomic library was constructed from extracted high molecular weight soil DNA as previously described in Hjort et al. (2010) and Hårdeman and Sjöling (2007).

Screening of the metagenomic library for chitinase genes

Clones of the fosmid library, i.e., 7,800 fosmids, were replica plated into 150 $\mu$L of Luria-Bertani (LB) medium supplemented with 12.5 $\mu$g/mL chloramphenicol and cultured over night at 37 °C. The clones were induced into high copy number with Fosmid Induction Solution (Epigenet, Nordic Biolabs, Täby, Sweden) for 5 h at 37 °C. For screening, aliquots from the 96-clone cultures of every microtitre plate were pooled, making a total of 78 superpools, representing the whole metagenomic library. Fosmid DNA was extracted from 150 $\mu$L of the 78 superpool cultures with R.E.A.L. Prep 96 Plasmid Kit (Qiagen, Hilden, Germany), following the manufacturer's instructions.

The metagenomic library, i.e., the super pools, was screened for family 18 chitinase genes by PCR using degenerate primers ChiA_F2 (5'-CGT GGA CAT CGA CTG GGA RTW YCC-3') and ChiA_R2 (5'-CCC AGG CGC CGT AGA RRT CRT ARS WCA-3')(Hobel et al. 2005), designed on the
basis of the conserved regions of the catalytic domain of glycoside hydrolase family 18. The PCR reactions were set up according to Hobel et al. (2004) with a few modifications as described in Hjort et al. (2010). Depooling was performed with a second round of PCR screening to identify the single fosmid clone containing a chitinase gene. For activity-based screening, super pools were analyzed for chitinolytic activity using the activity assay with 4-methylumbelliferyl-N-N-chitobiose as substrate, as described below.

Sequence analysis of the chi18H8 gene

The complete gene sequence of the identified chitinase chi18H8 was obtained by primer walking technique (Macrogen Inc., Seoul, Korea) from the primers described above, towards the downstream and upstream insert/fosmid junctions. A putative signal peptide in the Chi18H8 amino acid sequence was identified by SignalP 4.0 server (Petersen et al. 2011) and InterProScan 4.8 (Zdobnov and Apweiler 2001). Sequence similarity comparisons were performed with InterProScan and Prosite (Sigrist et al. 2010) softwares. Phylogenetic comparison by amino acid sequence alignment of the identified putative chitinase with homologous chitinases was performed with protein BLAST (Altschul et al. 1990). A phylogenetic tree using PhyML was constructed based on amino acid sequence alignments of the catalytic domain of Chi18H8 with that of representative of the eight groups of bacterial glycosyl hydrolases according to Karlsson and Stenlid (2009).

Subcloning of the chi18H8 gene

The chi18H8 gene was cloned into the expression vector pGEX-6P-3 (GE Healthcare, Upsala, Sweden), by first amplifying the gene with the primers Chi_18H8_F (5'-GGG CCC G AAT TCC ATG CGC CAG CTC ACG CTT CTC and Chi_18H8_R (5'-GGG CGC CTC GAG CTA TCA ATT GGC CCT ATG CAG ACT, with the positive fosmid clone as template. For the correct orientation of the gene into the vector, restriction enzyme sites EcoRI and XhoI were introduced, respectively, into the primers Chi_18H8_F and Chi_18H8_R, as underlined. A standard PCR reaction (Sambrook and Russell 2001) at 68 °C annealing temperature was performed with High Fidelity PCR Enzyme (Fermentas, Thermo Fisher Scientific Inc, VWR International, Stockholm, Sweden). The amplified fragment, double digested with EcoRI and XhoI was then ligated with T4-DNA ligase (New England Biolabs, Ipswich, England) into the expression vector pGEX-6P-3 (GE Healthcare) treated with EcoRI and XhoI and including the protein tag glutathione S transferase (GST). The resulting plasmid, pGEX-6P-3::chi18H8 was transformed into Escherichia coli TOP-10 (Invitrogen Life Technologies, Stockholm, Sweden) and sequenced to make sure that the gene was correctly inserted (Macrogen Inc). For overexpression, the construct pGEX-6P-3::chi18H8 was transformed into E. coli BL21 (DE3) cells (Invitrogen).

Expression and purification of the Chi18H8

For GST-Chi18H8 fusion protein overexpression conditions in E. coli BL21 (DE3), see the Electronic supplementary material. For protein purification, a single colony of E. coli pGEX-6P-3::chi18H8 was inoculated into 5 mL of LB medium with 50 mg/L ampicillin (Sigma-Aldrich, St. Louis, MO, USA) and incubated at 37 °C for 2 h at 200 rpm. The culture was inoculated into 300 mL Erlenmeyer flasks 50 mL malt extract (ME) medium (6 g/L malt extract, 1.8 g/L maltose, 6 g/L dextrose, and 1.2 g/L yeast extract: all medium components from Sigma-Aldrich) supplemented with 50 mg/L of ampicillin and incubated overnight at 25 °C and 150 rpm; 25 mL of the overnight culture was then inoculated into 2 L flasks with 375 mL ME medium and 25 mg/L ampicillin and incubated at 20 °C and at 150 rpm. At an O.D. of 0.4, isopropyl-β-D-thiogalactopyranoside (IPTG, Sigma-Aldrich) was added to a final concentration of 0.1 mM and the incubation continued for 48 h at 16 °C. Cells were harvested by centrifugation at 8,000×g for 15 min and then sonicated (6 cycles of 30 s each, with a 1 min interval using a Branson Sonifier 250, Danbury, USA) in phosphate-buffered saline (PBS; 2 mL/g cells) at pH 7.4 (140 mM NaCl, 2.7 mM KCl, 10 mM Na3HPO4, and 1.8 mM KH2PO4), containing 1 mM phenylmethylsulfonylfluoride (Sigma-Aldrich), 0.7 μg/mL of pepstatin A (Sigma-Aldrich) and 10 μg/mL deoxyribonuclease I (Sigma-Aldrich). The soluble fraction was separated from the insoluble fraction by centrifugation at 18,000×g for 1 h at 4 °C. The GST-Chi18H8 fusion protein was purified from the soluble fraction by loading on a GSTrap FF column 1 mL (GE Healthcare) using PBS at pH 7.4. For the on-column cleavage of the GST tag, PreScission buffer (50 mM TrisHCl at pH 7.0, 0.15 M NaCl, 1 mM ethylenediaminetetraacetic acid, and 1 mM diithiothreitol) containing 30 U of PreScission protease (GE Healthcare) was loaded. The column was then incubated for 15 h at 4 °C, followed by 2 h at room temperature. The cleaved Chi18H8 was eluted by PreScission buffer. The GST tag and the PreScission protease bound to the column were eluted by 50 mM TrisHCl at pH 8.0 containing 15 mM reduced l-glutathione, according to the manufacturer's protocol.

Protein and zymogram analyses

Protein samples from soluble and insoluble E. coli cell fractions and from fractions collected during purification were loaded onto 10 % (w/v) sodium dodecyl sulfate (SDS)–polyacrylamide gel for electrophoresis (PAGE). For the insoluble fractions, pellets obtained after sonication and centrifugation of E. coli cells, were resuspended in an appropriate volume of
Laemmli sample buffer. Protein bands were visualized in gels by staining with Coomassie brilliant blue (Laemmli 1970). For molecular mass determination, PageRuler™ Prestained Protein Ladder (Thermo Scientific, Milan, Italy) was used. Protein concentration was estimated by densitometric analysis of SDS gel bands using the Quantity One analysis software (Bio Rad, Hercules, CA, USA) and by the use of the extinction coefficient at 280 nm (77,015 M⁻¹ cm⁻¹) determined by urea denaturation compared with the theoretical extinction coefficient based on the amino acid sequence.

Zymogram was used to detect chitinolytic activity, using 10 % (v/v) polyacrylamide gel containing 0.7 mg/mL carboxymethyl-chitin–Remazol brilliant violet (CM-chitin–RBV; Loewe Biochemica, Sauerlach, Germany). The purified Chi18H8 was diluted in an appropriate sample buffer volume lacking any reducing agent and incubated for 10 min at room temperature. Electrophoresis was conducted at 4 °C in Tris–glycine–SDS running buffer according to the standard running conditions. The gel was then rinsed twice in 2.5 % (v/v) Triton X-100 for 30 min at room temperature to remove SDS and equilibrated in 0.1 M sodium acetate buffer (pH 5) at 37 °C. Clear zones indicating chitinolytic activity appeared after overnight incubation at 37 °C.

Chitinase activity assay

Enzymatic activity of the metagenomic library, fosmid clone crude extract, and purified enzyme was assayed fluorometrically by measuring the release of 4-methylumbelliferone (MU) from the chitooligosaccharide analogs 4-methylumbelliferyl-N-N′-N''-chitotriose (MU-NAG3), 4-methylumbelliferyl-N-N′-chitobiose (MU-NAG2), and 4-methylumbelliferyl-N-aceetyl-d-glucosamine (MU-NAG). In the case of metagenome library screening of the pooled fosmid clones, 100 μL of cell suspensions previously grown in LB medium supplemented with 12.5 μg/mL chloramphenicol, and induced by Fosmid Induction Solution (Epicentre) according to the manufacturer’s protocol, were added in black microtitre plates with 10 μL 10-mM MU-NAG2. The plates were incubated at 20 °C for up to 7 days. Fluorescence was monitored daily using a FLUOstar (BMG LABTECH, Offenburg, Germany) plate reader with an excitation wavelength at 320 nm and an emission at 460 nm. For assaying enzyme extracts, the standard reaction mixture (50 μL) contained: 0.4 mM of the chitooligosaccharide in 50 mM sodium acetate buffer pH 5 and 5 μL of protein sample. The reaction was stopped after 30 min at 37 °C by the addition of 150 μL of 400 mM sodium carbonate solution. Fluorescence was monitored as described above. All measurements were performed at least in triplicate. One unit of enzyme activity was defined as the amount of enzyme that hydrolyses 1 μmole of MU of the substrate per minute at pH 5.0 and 37 °C (McCreath and Gooday 1992; Di Maro et al. 2010).

The optimum pH for Chi18H8 activity was determined using the fluorescent assay described above with the following buffers (50 mM) at respective pH: sodium acetate (pH 4.0 and 5.0), sodium phosphate (pH 6.0, 7.0, and 8.0), and glycine–sodium hydroxide (pH 9.0) (Pears 1980). The optimum temperature for Chi18H8 activity was determined by the fluorescent assay incubating the reaction mixture at various temperatures (5–70 °C).

Antifungal activity assay

Antifungal activity of Chi18H8 was estimated by a growth inhibition assay on a set of eight plant pathogenic fungi, listed in Table 1. An agar plug, harboring an actively growing fungal culture, was placed onto the middle of a ME agar plate, pH 5, supplemented with 50 mg/L ampicillin and 0.1 mM IPTG. Subsequently, cell suspensions of E. coli BL21(DE3) pGEX-

| Fungal strain          | General crop disease and damage                        | Reference                  | Inhibited by Chi18H8 |
|------------------------|--------------------------------------------------------|----------------------------|----------------------|
| Alternaria alternata   | Leaf spot and other diseases on over 380 host species  | Mukherjee and Sen (2006) and Di Maro (2010) | +                     |
| Aspergillus niger      | Black mold on a certain fruits and vegetables, such as grapes, onions, and peanuts | Xiao et al. (2009), Yan et al. (2008), and Singh et al. (2007) | −                     |
| Colletotrichum gloeosporioides | Anthracnose disease on various temperate, subtropical, and tropical fruit trees worldwide | Prapagdee et al. (2008) and Singh et al. (2007) | +                     |
| Fusarium graminearum   | Head blight in oats and barley, resulting in yield reductions and contaminations of grains with trichothecene mycotoxins such as deoxynivalenol | Dehestani et al. (2010) | +                     |
| Fusarium oxysporum     | Has a broad host range. Individual isolates usually cause disease only on a narrow range of plant species | Wang et al. (2009) and Lee et al. (2007) | +                     |
| Penicillium chrysogenum | Post harvest pathogen of fruits and vegetables as it is widely distributed in nature | Mukherjee and Sen (2006) | −                     |
| Rhizopus stolonifer    | Bread mold fungus                                      | Yan et al. (2008)          | −                     |
| Trichoderma harzianum  | Major source of crop loss for mushrooms farmers        | Hassan et al. (2009)       | −                     |
The nucleotide sequence of the chitinase gene from the expression plasmid pGEX-6P-3::chi18H8 was induced by the IPTG added into the agar media. E. coli BL21(DE3) pGEX-6P-3 was used as negative control. The plates were incubated at 25 °C and growth of fungal hyphae was monitored using a microscope. The following fungal strains were all generously obtained as a gift from Prof. J. Schnürer’s research group, Department of Microbiology, Agricultural University of Sweden: Colletotrichum gloeosporioides CR21, Fusarium graminearum IBT 1958 (from Center of Microbial Biotechnology, Department of Systems Biology, Technical University of Denmark), Fusarium oxysporum FRR3414 (from Commonwealth Scientific and Industrial Research Organization, Food and Nutritional Sciences, North Ryde Sydney, NSW, Australia), Penicillium chrysogenum ATCC9179, and Trichoderma harzianum CBS226.95. Furthermore, the following strains from Schnürer’s research group’s own fungal collection were used: Alternaria alternata J414, Aspergillus niger J8, and Rhizopus stolonifer J45.

Sequence database deposition

The nucleotide sequence of the chitinase chi18H8 gene was deposited in the GenBank database under the accession number KC763366.

Results

Identifying a novel chitinase in the metagenomic library

The fosmid library, previously generated from a field soil reported to be suppressive for fungal phytopathogens, was screened by combining activity screening with a “genetic sieving” utilizing degenerate PCR primers for the glycoside hydrolases of family 18 A. For this purpose, a pooling strategy was successfully applied which made screening of the library more efficient. Those clones expressing chitinolytic activity in the presence of the synthetic chitin surrogate substrate oligosaccharide MU-NAG2 were considered as positive hits.

Among the 78 superpools screened, four pools returned a PCR product of the expected size (270 nucleotides). Each clone within the four positive super pools was further analyzed individually and 25 positive clones were identified in a second PCR run. From one of the clones (plate 18 row H8) that showed both strong chitinase activity and PCR signal, we identified the complete open-reading frame of a novel chitinase by primer walking technique. The gene, named chi18H8, consists of 1,275 nucleotides encoding a protein of 425-amino acid residues with a predicted molecular weight of 46 kDa. In silico sequence analysis identified a protein motif for a signal peptide of 20 amino acid residues at the N-terminal, followed by a putative catalytic domain (amino acid residues, 37 to 408) (Fig. 1a). The catalytic domain includes the consensus motif for the active site consisting of the amino acid residues FDGIDIDWE (amino acid residues, 175 to 183) with the motif DxDxE conserved among all the family 18 chitinase members. It also contains the so-called chitinase insertion domain (CID) between amino acid residues 310 and 379, identified by the conserved residues YxR. Chitin has been suggested to insert at this site to reach the active site, and it consists of a small α+β domain which is the main structural feature of subfamily A of family 18 chitinases (Li and Greene 2010).

A phylogenetic tree based on sequence alignments of the catalytic domain of Chi18H8 with that of representatives of the eight groups of bacterial glycosyl hydrolases (according to Karlsson and Stenlid 2009), shows that Chi18H8 belongs to the group II of chitinase family 18. Chi18H8 clusters with representatives of divergent phyla, such as proteobacteria (Saccharophagus) and actinobacteria (Streptomyces) (Fig. 1b) Protein sequence comparison of the full-length protein sequence showed highest amino acid identity (45 %) to a chitinase from an uncultured bacterium (Uchiyama and Watanabe 2006). The closest match to a protein sequence from an identified organism was with the following myxobacteria: Chondromyces apiculatus (41 % identity), Corallococcus coralloides (40 % identity), and Myxococcus xanthus (39 % identity).

Purification of the novel chitinase Chi18H8

Heterologous expression of the chi18H8 gene, which was cloned into a pGEX-6P-3 vector in E. coli gave the fusion protein GST-Chi18H8 of an estimated molecular mass of 73 kDa (Fig. 2a) and with a pI of 6.39. This expression system was selected as we successfully used it previously for purifying metagenomically sourced lipases and esterases (Hårdeman and Sjöling 2007) and others have used it for expressing bacterial chitinases (Tsujibu et al. 2000). The GST moiety of the vector adds at the amino terminus of the recombinant protein to facilitate on column purification via affinity chromatography as the GST-tag binds to the column and then can be cleaved off using a PreScission protease, allowing final recovery of the native protein (Makrides 1996).

Results from experiments conducted on heterologous fusion protein production by varying different cultivation conditions are reported in the Electronic supplementary material and shown in Figs. S1 and S2. In these experiments, enzymatic activity assay (i.e., the fluorimetric assay with MU-NAG ana-logs as substrates, Fig. S1) and detection of protein accumulation by SDS-PAGE (Fig. S2) revealed that the fusion protein maintained the chitinolytic activity, but most of it accumulated into insoluble fractions, from which protein recovery is known to be challenging (Huang et al. 2012). To facilitate purification, different expression trials (Electronic supplementary material) were conducted to increase the protein production in the soluble...
form. At the most optimal conditions we achieved so far (inducing cells growing in ME at early exponential phase, with 0.1 mM IPTG and harvesting the cells after 48 h at 16 °C), only 10 % of the chitobiosidase activity initially detected in the soluble fraction (34 U) was recovered in the purified fraction, starting from 2.3 g of cell paste. After the on-column PreScission protease cleavage, 30 μg high-purity Chi18H8 per liter of culture (21 μg/g cells) was obtained (Fig. 2a).

Enzyme characterization of Chi18H8

Notwithstanding the limited amount of the purified enzyme, characterization of the Chi18H8 protein was possible. Zymogram on carboxymethyl chitin confirmed the chitinolytic activity of the purified Chi18H8 (Fig. 2b), both with (lane 1) and without (lane 2) the GST-tag. Substrate specificity analyses of Chi18H8 against three different-length analogs of chitooligosaccharides showed an activity of 113 U/mg protein for MU-NAG2 as substrate, and of 39 U/mg protein on MU-NAG3, but no activity against MU-NAG. These substrates are commonly used for detecting chitobiosidase, endochitinase, and N-acetylglucosaminidase activities, respectively (Howard et al. 2003). Our results indicated that the Chi18H8 enzyme has a prevalent chitobiosidase activity.

The optimum temperature for Chi18H8 activity with both MU-NAG2 (Fig. 3a) and MU-NAG3 was 35 °C. The enzymatic activity was also tested at different pH ranging from 3 to 9. With a pH optimum between 3 and 5, more than 80 % of the
Chi18H8 activity remained at acidic and sub-acidic conditions. At pH above 6, the hydrolytic activity drastically decreased (Fig. 3b).

Chi18H8 has antifungal activity

The antifungal activity of Chi18H8 was determined by co-cultivating E. coli pGEX-6P-3::chi18H8 or pGEX-6P-3 strains, with the phytopathogen fungi listed in Table 1. The recombinant bacterial growth, surrounding the fungal plug, inhibited the growth of C. gloeosporioides, F. graminearum, and F. oxysporum as shown in Fig. 4, and A. alternata. No growth inhibition was observed in the negative control with the bacterial suspension harboring the plasmid without chi18H8.

Discussion

Access to the functional capacities of the bacterial community without limits of cultivation is important in the discovery of novel biocatalysts, not the least when the demand for more environmentally friendly alternatives is strong. We used a genetic and functional metagenomics approach to identify a novel chitinase with antifungal activity from a Swedish field soil, previously characterized as suppressive toward phytopathogens.

The novel chitinase Chi18H8 only shows 45 %, or less, amino acid sequence identity to any known chitinase but contains the conserved consensus sequences of family 18 chitinases including the conserved active site motif, DxDxE of the catalytic domain. Consistently with what has been shown for the catalytic domain of the family 18 chitinases (Suzuki et al. 2002), structure prediction showed that Chi18H8 has a triosephosphate isomerase fold (TIM barrel)
Chi18H8 over the trimer substrate analog MUF-NAG3, but it is Chi18H8 has a preference for the dimer substrate MUF-

Substrate binding cleft (Horn et al. 2006). Chitinases, such as Serratia marcescens ChiA and ChiB, are classified as exochitinases (endochitinase activity (Suzuki et al. 2002). Exochitinases are suggested to hydrolyze off oligosaccharide dimers or chitobiosidase from the polysaccharide by

According to studies on ChiA in S. marcescens (Zees et al. 2009), the presence of the CID is suggested to enhance the chitobiosidase activity as well as the processivity during degradation of the polysaccharide chains, facilitating the accessibility to crystalline and insoluble substrate, such as chitin.

The metagenome-sourced Chi18H8 shows optimal activity and stability at acidic pH and mesophilic temperatures, consistently with what has been reported for family 18 chitinases isolated from cultured microorganisms (Bhattacharya et al. 2007). Unlike other family 18 chitinases, however, the Chi18H8 seems to exert antagonistic selective activity against four important plant pathogens, coherently with the suppressive nature of the soil from which it was isolated. To date, antifungal activity has been reported among the family 19 of glycosyl hydrolases (Gherbawy et al. 2012). Family 19 chitinases differ from those belonging to family 18 in amino acid sequence, three-dimensional protein structure, molecular mechanism of catalytic region, and enzymology and are thus considered to have different evolutionary origin (Kawase et al. 2006). Even if further investigation should be conducted on the antifungal spectrum of the Chi18H8 by using the pure protein product, to our knowledge, this is the first report of a family 18 chitinase possessing an antifungal activity.

Further investigations on the mechanism of action and on the antifungal activity of Chi18H8 will be possible by obtaining a higher yield of the pure protein though improving the host-expression system. Low purification yield is a persistent bottleneck in heterologous expression in model hosts of metagenomically derived genes of unknown origin (Schmeisser et al. 2007; Sjöling et al. 2007). Limitations in availability of suitable alternative host systems have so far made E. coli based expression the most common for metagenomic screening and as a result most of the metabolic diversity present in environmental samples is less likely to be efficiently accessed. Success of protein expression depends on the host’s ability to transcribe the metagenomic DNA efficiently, translate the mRNA to form a functional protein, fold, and localize the protein correctly. In our case, detection of Chi18H8 in the E. coli hosted metagenomic library was facilitated by the high sensitivity of the chitobiosidase fluorimetric assay probably in combination with the occurrence of cell lysis during the incubation period. Indeed, Chi18H8 purification from E. coli was hampered by its accumulation within the insoluble fraction, i.e., inclusion bodies and /or membranes. Our recent trials (Berini et al., unpublished results) on cloning the chi18H8 gene into another E. coli vector (pET24b+, Novagen, Milan, Italy) providing only a histidine hexamer at the C terminus of the protein product (C-His6-Chi18H8), indicate that most of the protein accumulates into
the insoluble fraction as in case of the GST-Chi18H8 fusion protein. Thus, we could exclude that protein accumulation is caused by the length and the position of the GST-tag in the pGEX-6P-3 expression system. However, the chitinase was expressed in both systems with its putative signal peptide, which may, through its hydrophobic nature, facilitate aggregation into inclusion bodies and/or sticking to E. coli membranes. A previous paper from other authors (Hoster et al. 2005) reported that a tenfold higher heterologous production of a Streptomyces chitinase in E. coli was achieved when the gene was cloned without the region encoding the putative signal sequence. Alternatively, the development of solubilisation and refolding protocols (Huang et al. 2012) or the use of expression systems alternative to E. coli are being investigated to allow larger scale production of Chi18H8 required for further detailed protein characterisation and for its in-field evaluation as a novel biocontrol agent.

To our knowledge, this is the first active chitinase identified, produced, and purified using a metagenomic approach. This study confirms that accessing the metabolic functions of a suppressive soil by functional metagenomics can provide a novel chitinase with antifungal activity that has the potential application as a biocontrol compound against crop fungal pathogens and may offer a more environmentally sustainable alternative to the otherwise often toxic chemical fungicides.

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