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#### Postharvest application of a novel chitinase cloned from Metschnikowia fructicola and overexpressed in Pichia pastoris to control brown rot of peaches

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1	Postharvest application of a novel chitinase cloned from Metschnikowia fructicola and
2	overexpressed in Pichia pastoris to control brown rot of peaches
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4	Short running head:
5	Chitinase against brown rot of peaches
6	
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#### 22 Abstract

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Metschnikowia fructicola strain AP47 is a yeast antagonist against postharvest pathogens of fruits. 24 The yeast was able to produce chitinase enzymes in the presence of pathogen cell wall. A novel 25 chitinase gene MfChi (GenBank accession number HQ113461) was amplified from the genomic 26 DNA of *M. fructicola* AP47. Sequence analysis showed lack of introns, an open reading frame (ORF) 27 28 of 1,098 bp encoding a 365 amino acid protein with a calculated molecular weight of 40.9 kDa and a predicted pI of 5.27. MfChi was highly induced in Metschnikowia fructicola after interaction with 29 Monilinia fructicola cell wall, suggesting a primary role of MfChi chitinase in the antagonistic 30 31 activity of the yeast. The *MfChi* gene overexpressed in the heterologous expression system of *Pichia* pastoris KM71 and the recombinant chitinase showed high endochitinase activity towards 4-32 Nitrophenyl β-D-N, N', N''-triacetylchitotriose substrate. The antifungal activity of the recombinant 33 34 chitinase was investigated against Monilinia fructicola and Monilinia laxa in vitro and on peaches. The chitinase significantly controlled the spore germination and the germ tube length of the tested 35 pathogens in PDB medium and the mycelium diameter in PDA. The enzyme, when applied on 36 peaches cv. Redhaven, successfully reduced brown rot severity. This work shows that the chitinase 37 MfChi could be developed as a postharvest treatment with antimicrobial activity for fruit undergoing 38 39 a short shelf life, and confirms that *Pichia pastoris* KM71 is a suitable microorganism for costeffective large-scale production of recombinant chitinases. 40

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42 Keywords: Cloning, Quantitative real-time PCR (RT-qPCR), recombinant expression, *Pichia*43 *pastoris, Monilinia fructicola, Monilinia laxa*

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Considerable losses are caused by postharvest diseases during transportation and storage of fruit 50 (Sharma et al., 2009). Brown rot caused mainly by Monilinia laxa (Aderh. et Rulh.) Honey and 51 Monilinia fructicola (G. Wint.) Honey is considered the main postharvest disease of stone fruit (De 52 Cal and Melgarejo, 1999; De Cal et al., 2009). M. fructicola is the most destructive pre- and 53 54 postharvest pathogen in all stone fruit-growing regions of the United States (Janisiewicz et al., 2013). In the European Union, neither additional cultural measures nor increased fungicide treatments are 55 sufficient to control brown rot in the orchard and in postharvest after the introduction of M. fructicola 56 57 (EFSA, 2011; Pellegrino et al., 2009), and no chemical fungicides are allowed for postharvest treatment of stone fruit. Moreover, the public demands to reduce pesticide use on fruit and to improve 58 environmental protection and human health have increased the need to develop alternative control 59 60 methods (Lopez-Reyes et al., 2013; Sisquella et al., 2014). Biological control using antagonistic yeasts has been explored as one of several promising alternatives to chemical fungicides (Liu et al., 61 62 2013a). Antagonistic yeasts deserve particular attention and are considered promising biocontrol candidates, as their activity neither involves production of toxic metabolites nor negative impact on 63 64 the environmental safety (Spadaro et al., 2002; 2008).

65 Among different antagonistic yeasts, Metschnikowia fructicola Kurtzman and Droby is an important yeast species which has been successfully applied to control a number of pathogens on fruits and 66 vegetables, such as *Penicillium expansum* on apple (Liu et al., 2011; Spadaro et al., 2013), *Botrytis* 67 68 cinerea on grape (Karabulut et al., 2003; Kurtzman and Droby, 2001) and on strawberries (Karabulut et al., 2004). Moreover, one strain of Metschnikowia fructicola was registered and commercially 69 70 available in Israel to control storage diseases of fruits and vegetables (Kurtzman and Droby, 2001, 71 Macarisin et al., 2010). The strain AP47 of Metschnikowia fructicola (Zhang et al., 2010a) was obtained from the carposphere of an apple grown in organic orchard in North Italy. Under semi-72 commercial conditions, Metschnikowia fructicola strain AP47 showed a high efficacy in controlling 73

brown rot caused by *Monilinia* spp. on stone fruits, however its mechanism against postharvest
pathogens is still unclear (Zhang et al., 2010a).

Various mechanisms of action of antagonistic yeasts have been described, such as competition for nutrients and niche exclusion (Li et al., 2008; Liu et al., 2012a), induction of host defense mechanisms (Jiang et al., 2009; Xu et al., 2013) and the production of hydrolases such as chitinase, protease and glucanase, which is proposed as an important mode of action against fungal pathogens, due to its role in breaking down pathogens cell wall and inhibiting spore germinations (Masih and Paul, 2002; Smits et al., 2001; Zhang et al., 2011; 2012).

Cloning, expression and characterisation of new chitinase genes from microorganisms is useful for antagonism activity as well as for developing new potential chitin biological degraders. Compared with the extensive research into the chitinases from some antagonistic fungi, such as *Trichoderma* spp. (Nakahara et al., 2001; Silva et al., 2011) and bacteria such as *Bacillus* spp. (Shivakumar et al., 2014; Yang et al., 2009), few studies have been carried out on chitinases produced by yeasts with molecular tools. To our knowledge, there is no published report on cloning and phylogenetic analysis and expression of chitinase from the antagonistic yeast species *Metschnikowia fructicola*.

Recently *Pichia pastoris* has emerged as an important yeast host for heterologous protein expression
(Cregg et al., 1993; Macauley et al., 2005), since it has many of the advantages of higher eukaryotic
expression systems, such as protein processing and folding and posttranslational modifications
(Balamurugan et al., 2007). Therefore it was used in this study for chitinase expression.

The objectives of this research were: i) to study the chitinolytic activity of the antagonistic yeast *Metschnikowia fructicola* strain AP47 *in vitro*; ii) to clone and characterize the chitinase gene *MfChi* from AP47; iii) to analyse *MfChi* gene expression in AP47 after exposure to pathogen cell wall preparation through reverse transcription quantitative PCR (RT-qPCR); iv) to express the chitinase MfChi in the methylotrophic yeast *Pichia pastoris;* v) to study the antifungal activity of the expressed chitinase *in vitro* and *in vivo* and the effect of the enzyme concentration on the control of *M. laxa* and

99 *M. fructicola*.

#### 101 **2.** Materials and Methods

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#### 2.1. Microorganisms, growth media, plasmids and molecular kits

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Metschnikowia fructicola Kurtzman and Droby strain AP47 (Zhang et al., 2010a) was isolated from 105 106 the carposphere of apple cv. Golden delicious, harvested in an organic orchard located in Piedmont, 107 Northern Italy and identified by using molecular and morphological tools. The microorganism culture was stored at -80 °C in cell suspension with 65% (v/v) glycerol and 35% (v/v) of a solution of 100 108 109 mM MgSO<sub>4</sub> and 25 mM Tris (pH 8.0). Yeast subcultures were grown in YEMS (30 g/L yeast extract, 5 g/L D-mannitol, 5 g/L l-sorbose (Spadaro et al., 2010). Five strains of Monilinia fructicola (G. 110 Wint.) Honey and five strains of Monilinia laxa (Aderhold & Ruhland) Honey isolated from rotted 111 112 peaches were used as a mixture throughout this work after being selected for their virulence by inoculation in artificially wounded peaches. 113

114 Oligonucleotides, pGEM-T vector and Escherichia coli strain JM109 used in this study were purchased from Promega (Madison, WI, USA). The kits of DNA and RNA extraction (DNeasy and 115 RNeasy). QIAquik PCR purification, Reverse-transcript PCR, Plasmid-extraction, QIAquik Gel 116 117 extraction and one step RT-PCR kit as well as the materials for PCR were purchased from Qiagen (Hilden, Germany). The kit "Gene Walking Made Easy" and other materials for enzyme assays were 118 purchased from Sigma-Aldrich (St. Louis, MO,USA). Pichia pastoris KM71 strain used as host for 119 transformations with the plasmid pPIC9 and Escherichia coli strain DH5a used as host for the 120 plasmids were obtained from Invitrogen (Life Technologies, Carlsbad, USA). TURBO DNase was 121 purchased from Ambion (Ambion, Foster City, CA, USA). iScript cDNA Synthesis Kit and 2× Power 122 syber green supermix were purchased from Bio-Rad (Richmond, CA, USA) for RT-qPCR. 123

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### 125 2.2. Chitinase activity of the strain AP47 grown in vitro

127 To study the chitinase enzyme production from the strain AP47, and the effect of different substrates on its chitinolytic activity, the yeast strain was cultured in modified Lilly-Barnett minimal salt 128 (LBMS) medium (Lilly and Barnett, 1951) containing 2 mg/mL Monilinia fructicola cell wall 129 preparation (CWP), glucose or 5 mg/mL colloidal chitin as sole carbon source. CWP of the pathogen 130 Monilinia fructicola was prepared as described by Saligkarias et al., (2002), and colloidal chitin was 131 132 prepared according to the method described by Roberts and Selitrennikoff (1988) from shrimp shell chitin (C9752, Sigma-Aldrich). In preliminary experiments, the yeast strain produced the highest 133 chitinase activity when grown for 48 h. Therefore, we just measured the chitinase activity of the strain 134 135 when grown for 48 h. The spectrophotometric assay of chitinase activity was carried out according to the procedure developed by Miller (1959), with small modifications. Chitinase activity was 136 determined colorimetrically by using colloidal chitin as substrate. The reaction mixture, consisting of 137 138 500 µL colloidal chitin (0.5% w/v) and 500 µL enzyme solution, was incubated at 50 °C in a water bath for 30 min. The reaction was stopped by centrifugation at  $3,000 \times g$  for 3 min. An aliquot of the 139 140 supernatant (0.8 mL) was pipetted into a new sterile tube followed by adding 500 µL dinitrosalicylic acid. The reaction mixture was immediately boiled for 5 min. After cooling, the reducing sugars 141 released as chitinase activity were measured at 540 nm. One unit of chitinase activity was defined as 142 143 the amount of enzyme which produced 1  $\mu$ M/min reducing *N*-acetyl-D-glucosamine.

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#### 145 2.3. Cloning the chitinase gene MfChi from the genomic DNA

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The strain AP47 was grown in liquid medium YPD (20 g D-glucose, 20 g peptone casein, and 10 g yeast extract per litre) at 25 °C for 48 h, then centrifuged at 5,000×g for 10 min. DNA was extracted from the pellet with DNeasy extraction kit (Qiagen), according to the manufacturer's instruction.
To clone the first partial sequence of the chitinase gene from genomic DNA of *Metschnikowia fructicola* strain AP47, PCR amplification of the extracted DNA was performed by using the

degenerate primers 5'-CTNCTNTCNCTNGTNGTN-3' (Forward primer DPf) and 5'-152 CARTARTTRTTRTARAAYTG-3' (reverse primer DPr). DPf and DPr were designed according to 153 the conserved protein sequences (LLSLGG and QFYNNYC) obtained with DNAMAN 7.0 by using 154 the alignment of the deduced amino acid sequences of 8 yeasts chitinase genes deposited (Suppl. Fig. 155 1). After loading on agarose gel, PCR products were purified with QIAquick gel extraction kit 156 (Qiagen) according to the supplier's instructions, then ligated into pGEM-T cloning vector 157 (Promega), followed by transformation into chemically competent cells of E. coli strain DH5a 158 (Invitrogen) and selection of positive transformants with blue / white screening technique. The 159 sequencing and BLAST analysis showed that a fragment of 350 bp was obtained. To amplify and 160 identify the 5' and 3' flanking regions of the chitinase gene from the genomic DNA of Metschnikowia 161 fructicola strain AP47, special restriction digestion enzymes and primers were designed according to 162 the obtained sequence and the kit "Gene Walking Made Easy" (UVS1, Sigma-Aldrich, USA): AP47-163 164 5UTR: 5'-TCAGTCAAGAACGACAAGATCACAGTGTCC-3' and AP47-3UTR: 5'-TGATATGGACAAGAAGAAGCCTTTTGACTTGAACAAG-3' together with Vectorette Cla I 165 library of "genomic walking kit". The specific process was performed according to the supplier's 166 instructions. The fragment from Vectorette Cla I library of the strain AP47 was purified, ligated into 167 pGEM-T cloning vector and sequenced as described above. Finally the whole sequence of the targeted 168 gene was assembled, designated as MfChi and deposited in GenBank (accession number: 169 HQ113461.1). 170

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# 172 *2.4. Cloning the chitinase gene MfChi from the cDNA*

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Total RNA was extracted from the strain AP47 grown for 48 h in YPD broth at 25 °C by using an RNeasy Mini Kit (Qiagen), then treated with TURBO DNase (Ambion) according to the manufacturer's instructions. The absence of genomic DNA contamination was confirmed by PCR amplification of the housekeeping gene Actin1 (Li et al., 2006) using One Step RT-PCR Kit (Qiagen).

First-strand cDNA was synthesized using iScript cDNA Synthesis Kit (Bio-Rad) according to the 178 manufacturer's instructions. To amplify the chitinase gene MfChi from the cDNA of Metschnikowia 179 fructicola strain AP47, specific primers were designed according to the chitinase gene sequence 180 obtained from the genomic DNA of Metschnikowia fructicola strain AP47: forward primer (FP) 5'-181 ATGTTGATGCAACCATTTTTATGC -3' and primer (RP) 5'-182 reverse TCAGACTTTGAACTTTGGCTTG-3', then PCR products were purified and sequenced as 183 184 described above.

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#### 186 2.5. Analysis of MfChi gene expression

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AP47 was cultured in LBMS medium containing 2 mg/mL of Monilinia fructicola CWP at 23 °C by 188 shaking at 100 rpm, then collected after 6 h, 12 h, 24 h and 48 h of incubation. AP47 grown without 189 190 CWP served as a control. Each treatment consisted of three replicates at each time point and the experiment was repeated three times. Total RNA and cDNA synthesis were performed as described 191 192 previously, then the resulting cDNA was used as a template for RT-qPCR to quantify the MfChi 193 transcript expression under different time points. RT-qPCR was performed in triplicate on the cDNA obtained from each biological replicate using the 2× Power syber green supermix (Bio-Rad) for the 194 reaction mix according to the manufacturer's instructions. Amplification and detection were carried 195 out in an iCycler (Bio-Rad), set up with initial denaturation at 95 °C for 10 min followed by 40 cycles 196 comprising a denaturation step at 95 °C for 15 s and an annealing step at 60 °C for 1 min. The primers 197 198 *MfChi*-F (5'-TGATTTCCCCAAGATGAAGC-3') and *MfChi*-R (5'-AAAGTCACGAGCCTCTGCAT-3') were designed to optimally amplify *MfChi* gene sequence, and 199 transcript levels of Actin1 served as an internal standard. The primers used were Act1 F (5'-200 CCTGAGGAACACCCAGTCTT-3') and Act1 R (5'-GAGTTGTAAGTGGTTTGGTCG-3') 201 according to Liu et al. (2011). The expression ratio was calculated from equation  $2^{-\Delta\Delta CT}$ , where  $\Delta\Delta CT$ 202

represents the  $\Delta CT_{sample}$ - $\Delta CT_{control}$  (Livak and Schmittgen, 2001), and values were normalized to the control at 6 h, arbitrarily set to unity.

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#### 206 2.6. *Heterologous expression of MfChi in Pichia pastoris*

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The chitinase gene MfChi was amplified from Metschnikowia fructicola cDNA by PCR using 208 forward primer (5'-TCAGAATTCATGTTGATGCAACCATTTTTATGC-3') and reverse primer 209 (5'-CAGGAATTCTCAGACTTTGAACTTTGGCTT-3'); bases underlined encode 210 *Eco*RI restriction sites. The resulting DNA fragment (1098 bp) was digested with EcoRI before being ligated 211 into the corresponding sites of the expression vector pPIC9 and designated as pPIC9-MfChi. The 212 ligation product was transformed into E. coli JM109 (Promega) and the plasmid was sequenced at 213 BMR Genomics (Padova, Italy). Transformation of MfChi gene into P. pastoris KM71 was performed 214 215 as recommended by the manufacturer (Invitrogen). Briefly, pPIC9-MfChi was linearized using Stu I then transformed into competent P. pastoris KM71 cells via electroporation. The empty vector pPIC9 216 217 was also transformed as a negative control. Finally His<sup>+</sup> transformants of *P. pastoris* KM71 were 218 purified on minimal medium plates without histidine to ensure pure clonal isolates.

Transformed *P. pastoris* isolates were grown in 100 mL of Buffered Complex Glycerol Medium (BMGY) until the culture reached an OD 600 nm of 2-6, then the pellet was resuspended in 20 mL of Buffered Complex Methanol Medium (BMMY). Methanol was added at every 24 h interval to a final concentration of 1% to maintain the induction.

To analyze expression levels and the optimal time post-induction for harvest, supernatants were collected at different time points (0, 24, 48, 72, 96, 120 and 144 h) and secreted proteins were analyzed by SDS-PAGE (Laemmli, 1970) (Amersham ECL Gel 10%, GE Healthcare Life Science, Uppsala, Sweden). The recombinant protein MfChi was purified following the method of Liu et al., (2013b), then protein concentration was determined according to Bradford (1976) by using bovine serum albumin (Sigma–Aldrich) as a standard.

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#### 2.7. Recombinant chitinase activity assay

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Recombinant MfChi chitinase activity was determined using a colorimetric Chitinase assay kit (CS0980, Sigma-Aldrich) following manufacturers instruction. The absorbance was measured at 405 nm, then the specific activity of chitinase was expressed as U/mg, where one unit will release 1.0 micromole of *p*-nitrophenol from the substrate at pH 4.8 and 37 °C in one minute for each milligram of protein. Supernatant from cell culture of transformed *P. pastoris* with empty vector pPIC9 was used as a negative control. Three replicates in each treatment were performed, and the experiment was repeated three times.

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#### 240 2.8. Effect on pathogen mycelium growth in vitro

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The activity of the recombinant chitinase MfChi against *Monilinia fructicola* was assayed in Petri dishes containing PDA according to Banani et al. (2014) with some modifications. In brief, the recombinant chitinase MfChi was streaked into a PDA plate. A *M. fructicola* mycelial plug (5 mm diameter) was corked from a PDA culture and fixed in Petri dish at the same distance from the enzyme streak and the Petri dish border (control). After 6 days of pathogen growth at 25 °C, mycelial inhibition was measured and direct interaction *in vitro* was observed using an optical microscope (Eclipse 55i, NIKON, Tokyo, Japan).

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#### 253 2.9. Effect on pathogen spore germination and germ tube elongation

The effect of the recombinant chitinase MfChi was tested on conidia germination of *M. fructicola* 255 and M. laxa using the method of Zhang et al. (2012) with some modifications. In brief, tubes 256 containing 2.4 mL potato dextrose broth medium (PDB, Merck), 300 µL of Monilinia spp. conidial 257 suspension ( $1 \times 10^6$  conidia/mL) and 300 µL of the recombinant chitinase were co-incubated at 25 °C 258 on a rotary shaker (200 rpm). Two chitinase concentrations were assayed: C1 (7 ng/µL) and C2 (70 259  $ng/\mu L$ ). The control treatment consisted of water added to the tubes instead of the enzyme solution. 260 After 9 h and 18 h of incubation, 100 conidia of Monilinia spp. per replicate were observed, and their 261 germination rate (%) and germ tube length ( $\mu$ m) were measured by using an optical microscope. For 262 each treatment, three replications of three tubes were performed and the experiment was repeated 263 three times. 264

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#### 2.10. *Efficacy against Monilinia spp. in vivo*

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The biocontrol activity of the recombinant chitinase in controlling M. fructicola and M. laxa on 268 peaches cv. Redhaven was evaluated using the method of Yan et al. (2008) with some modifications. 269 270 Three wounds (4 mm deep  $\times$  3 mm wide) were made at the equator of each fruit. 20  $\mu$ L of recombinant chitinase were applied into each wound. The yeast AP47 was applied at  $10^8$  cells /mL (20  $\mu$ L) in 271 order to compare its activity with the chitinase. Peaches inoculated with Monilinia spp. spore 272 suspension acted as untreated control and peaches inoculated and treated with 2.5 mL/L of 273 tebuconazole (Folicur, Bayer Crop Science, Monheim, Germany; a.i.: 25.0%) were the chemical 274 control. Two hours later, 20 µL of pathogen suspension (10<sup>5</sup> conidia/mL) was inoculated into each 275 wound. Two chitinase concentrations (C1: 7 ng/µL and C2: 70 ng/µL) were used. The treated fruits 276 were incubated at 23 °C, and the rot diameter was measured 3 and 5 days after inoculation (DAI). 277 Each treatment consisted of three replicates with ten fruits per replicate and the experiment was 278 performed three times. 279

281 2.11. DNA sequence and phylogenetic analysis of the chitinase gene

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283	BLAST and ORF Finder programs at the National Center for Biotechnology Information (NCBI)
284	were used for the nucleotide sequence analysis, the deduction of the amino acid sequence and
285	database searches. Multiple sequence alignments of DNA and amino acid sequence were performed
286	using the programs of DNAMAN 7.0 and CLASTALW. The phylogenetic tree of the chitinase gene
287	was generated by MEGA6 using neighbour-joining method.
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289	2.12. Statistical analysis
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291	All statistical analyses were performed with SPSS version 20.0 (SPSS Inc., Chicago, IL, USA). Data
292	obtained in all the experiments were analysed using analysis of variance (ANOVA). The treatment
293	means were separated at 5% significance level by using Duncan's multiple range tests. Values are
294	presented as the mean $\pm$ SD (standard deviation of the mean). The results are the mean of three
295	independent experiments.
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297	3. <b>Results</b>
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299	3.1. Production of chitinase by Metschnikowia fructicola AP47 and its activity
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301	The strain AP47 showed chitinase activity when grown in different media. The chitinase activity of
301 302	The strain AP47 showed chitinase activity when grown in different media. The chitinase activity of AP47 was higher when grown in LBMS with <i>Monilinia fructicola</i> CWP as sole carbon source (0.35
301 302 303	The strain AP47 showed chitinase activity when grown in different media. The chitinase activity of AP47 was higher when grown in LBMS with <i>Monilinia fructicola</i> CWP as sole carbon source (0.35 U/mL) than with glucose (0.21 U/mL). The highest chitinase activity (0.46 U/mL) was observed when
301 302 303 304	The strain AP47 showed chitinase activity when grown in different media. The chitinase activity of AP47 was higher when grown in LBMS with <i>Monilinia fructicola</i> CWP as sole carbon source (0.35 U/mL) than with glucose (0.21 U/mL). The highest chitinase activity (0.46 U/mL) was observed when grown in LBMS with colloidal chitin.
<ul> <li>301</li> <li>302</li> <li>303</li> <li>304</li> <li>305</li> </ul>	The strain AP47 showed chitinase activity when grown in different media. The chitinase activity of AP47 was higher when grown in LBMS with <i>Monilinia fructicola</i> CWP as sole carbon source (0.35 U/mL) than with glucose (0.21 U/mL). The highest chitinase activity (0.46 U/mL) was observed when grown in LBMS with colloidal chitin.

308 PCR amplification of the partial sequence of chitinase gene from Metschnikowia fructicola AP47 DNA with the degenerate fragments DPf and DPr produced a 350 bp fragment of the putative 309 chitinase gene containing the consensus motif (DGXDFXXE) as signature pattern of Family 18 310 hydrolases. The signature pattern is highly conserved among most known chitinases from bacteria 311 and yeasts in its deduced amino acid sequence. PCR amplification of the flanking regions of the 312 313 fragment of the chitinase gene was performed from the genomic DNA of AP47, then the whole chitinase gene sequence was assembled and designated as MfChi (GenBank accession number 314 HQ113461) with 1,098 bp (Suppl. Fig. 2). PCR amplification of the cDNA of Metschnikowia 315 316 fructicola strain AP47 and sequence analysis revealed that the ORF from the cDNA of Metschnikowia fructicola strain AP47 shared the same nucleotide sequence with that of the genomic DNA, showing 317 the lack of introns inside the gene *MfChi*. 318

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#### 320 *3.3. Characterization of the chitinase gene MfChi*

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322 The prediction of the signal peptide of the chitinase gene *MfChi* was performed according to Bendtsen et al. (2004), and showed the presence of 19 amino acid signal peptide (positions from 1 to 19). Often, 323 324 the first 20 amino acids serve as a typical cleavable signal sequence for secreted proteins (Kuranda and Robbins. 1991). Sequencing and nucleotide analysis confirmed the lack of introns inside MfChi 325 gene, and an ORF of 1,098 bp encoding a 365 amino acid protein with predicted molecular weight of 326 327 40.9 kDa and pI of 5.27 were calculated (Suppl. Table 1). Prediction and analysis of the deduced amino acids from the gene showed that MfChi has 14 putative phosphorylation sites at positions of 328 329 34, 49, 103, 108, 115, 233, 285, 296, 125, 288, 92, 159, 204 and 304, respectively. In addition, MfChi has 4 putative N-glycosylation sites at positions of 67, 314, 318 and 324, respectively (Suppl. Fig. 2), 330 while no O-glycosylation sites were observed. Sequence alignment of the deduced amino acids from 331 MfChi with other related chitinases of yeasts retrieved from NCBI database showed that MfChi has 332

only high similarity (97.0%) with MpChi chitinase of Metschnikowia pulcherrima strain MACH1 333 (GenBank accession number HQ113462, Saravanakumar et al., 2009), but low similarity to other 334 yeast chitinases (Suppl. Fig. 3). However, when alignment was performed with the N-terminal regions 335 336 of the chitinase MfChi and other yeast chitinases, a high homology was observed between these chitinases (Suppl. Fig. 4). Moreover MfChi shared a common putative catalytic domain which 337 conformed to the signature motif (DXXDXXXE) of family 18 of chitinases proposed by Watanabe 338 et al. (1993), suggesting to belong to family 18 of chitinases. In addition, six conserved cysteine 339 residues required for substrate-binding by the chitinase were identified (Suppl. Fig. 4). To reveal the 340 relationship of MfChi with the chitinases from other yeast and other organisms, a phylogenetic 341 analysis was performed on the nucleotide sequences. MfChi belong to GH family 18 and it is included 342 into subgroup II including yeast chitinases (Suppl. Fig. 5 and Suppl. Table 2). Among the chitinases 343 of the subgroup II, MfChi is the closest to the yeast chitinase MpChi of Metschnikowia pulcherrima 344 345 strain MACH1 (Saravanakumar et al., 2009).

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#### 347 *3.4. Expression of MfChi in Metschnikowia fructicola in response to pathogen cell wall*

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Quantitative real-time PCR (RT-qPCR) was conducted to analyze the expression of *MfChi* in AP47 after exposure to CWP of *Monilinia fructicola in vitro*. The time-points included in the analysis were 6 h, 12 h, 24 h and 48 h of co-incubation. The expression of the *MfChi* gene was upregulated at an early stage of incubation and then it was downregulated after 24 h of incubation. The results indicated that *Monilinia fructicola* CWP directly induced *MfChi* expression in *Metschnikowia fructicola*, especially at 12 hours of incubation, when the gene expression was threefold higher than without CWP (Fig. 1). At longer incubation times, *MfChi* expression gradually decreased.

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#### 357 *3.5. Expression and purification of recombinant chitinase MfChi*

The recombinant chitinase expressed in different *P. pastoris* isolates was analysed with SDS-PAGE (Fig. 2). After 120 h induction, a wide band appeared in some transformed isolates, with a size of about 40.9 kDa, which corresponds to the same molecular weight predicted, while no band was observed in the negative control (non-insert control: lane number 0). After small-scale production, the best producer colonies (isolate 2 and 4, Fig. 2) were selected for large-scale chitinase expression and purification.

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#### 3.6. Identification of MfChi enzyme activity

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Chitinase activity was performed to assess whether or not the expression of the chitinase MfChi in 368 the *P. pastoris* expression system resulted in a functional protein, and to evaluate its chitinolytic 369 activity. The recombinant chitinase MfChi showed high endochitinase activity towards the chitin 370 371 pseudosubstrate 4-Nitrophenyl β-D-N, N', N''-triacetylchitotriose p-(GlcNAc)3, which is a suitable substrate for endochitinase activity detection. Additionally, no chitinase activity was detected in the 372 373 culture medium of P. pastoris KM71 transformed with the empty vector pPIC9 after methanol 374 induction, indicating that chitinase displayed in the transformed yeast cells was due to the expression of the foreign gene MfChi. 375

376

## 377 *3.7. Effect on Monilinia spp. mycelium growth in vitro*

378

After 6 days of *Monilinia fructicola* growth in PDA plates streaked with the chitinase MfChi, the effect of the recombinant enzyme on pathogen mycelium growth was observed. MfChi chitinase significantly inhibited *M. fructicola* mycelial growth, in addition, no conidia sporulation was observed in the growing side of the pathogen mycelium closer to the chitinase treatment -.

383 This result was confirmed by observation under optical microscope which showed that the presence

of chitinase caused swelling of *M. fructicola* hyphae (Data not shown).

# 386 *3.8. Effect on Monilinia spp. spore germination and germ tube elongation*

387

By co-culturing both pathogens and the enzyme in liquid medium (PDB), the recombinant chitinase 388 MfChi efficiently controlled the conidial germination and germ tube length of *M. fructicola* and *M.* 389 *laxa* compared to the control treatments (pathogen+water) either at 9 h or at 18 h of incubation (Fig. 390 3). Moreover, the chitinase applied at higher concentration (C2: 70 ng/µL) showed better results in 391 392 reducing the spore germination (Fig. 3A) and the germ tube elongation, than applied at lower concentration (C1: 7 ng/ $\mu$ L). At 70 ng/ $\mu$ L, the chitinase almost completely blocked the germ tube 393 394 development of *M. fructicola* and *M. laxa* either at 9 h or 18 h of incubation (Fig. 3B). Higher germ tube lengths were observed for *M. fructicola* than for *M. laxa*, when incubated with water as control 395 (Fig. 3B). 396

397

#### 398 *3.9.Antifungal activity of recombinant chitinase against Monilinia spp. on peaches*

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400 The antifungal activity of the recombinant chitinase was investigated on peaches stored at room temperature. After 3 DAI (Fig. 4), the chitinase treatment significantly reduced Monilinia spp. rot 401 402 diameter compared to *M. fructicola* (33 mm) and (24 mm) *M. laxa* untreated controls. The chitinase at 70 ng/µL (C2) significantly controlled the lesion diameter on peaches (about 13 mm for both 403 pathogens), similarly to the antagonistic cells of Metschnikowia fructicola AP47 (10 mm), and better 404 405 than the protease at 7 ng/µL (C1) for *M. fructicola* (23 mm) and *M. laxa* (20 mm). At 5 DAI for *M.* fructicola, the chitinase was still more efficient than the untreated control (68 mm) especially at 70 406 ng/µL (50 mm), but its efficacy was lower than AP47 cell suspension (32 mm). For M. laxa (Fig. 4), 407 the rot diameter of peaches treated with the chitinase was similar to the untreated control. 408

409

#### 410 **4. Discussion**

This is the first study to characterize the role of a chitinase from *Metschnikowia fructicola* in the control of brown rot.. Though several studies have been performed by a large number of laboratories on the antagonistic activity of *Metschnikowia fructicola*, its mechanism of action against postharvest pathogens is still unclear, and the production of lytic enzymes, especially chitinases, is proposed as an important mode of action of antagonistic yeasts. Hydrolases attack the cell wall of phytopathogenic fungi to cause cell lysis and subsequent death (Tseng et al., 2008).

A better understanding of the modes of action of yeast biocontrol agents is essential for developing
appropriate commercial formulations and application methods, to maximize their potential use to
manage postharvest diseases (Droby et al., 2009; Zhang et al., 2010b).

In this research, the antagonistic yeast *Metschnikowia fructicola* strain AP47 showed to produce higher chitinase activity in the presence of *Monilinia fructicola* CWP and colloidal chitin, compared to glucose as sole carbon source in the medium.Therefore, we cloned,characterized, and expressed a novel endochitinase gene *MfChi* from *Metschnikowia fructicola* AP47 and we studied its antifungal activity and potential use against *Monilinia* spp. *in vitro* and *in vivo*.

The presence of a 19-residue putative signal peptide confirmed that MfChi is an extracellular protein,
a feature common to the majority of endochitinases expressed by mycoparasites (Hayes et al., 1994;
Morissette et al., 2003; Viterbo et al., 2001) and its activity towards p-(GlcNAc)3 substrate confirms
its endochitinase activity.

RT-qPCR expression analysis clarified that *MfChi* gene of *Metschnikowia fructicola* is highly induced by cell wall fragments of *Monilinia fructicola* during the first 24 h of contact then the gene was downregulated. Similar results were reported for the endochitinase chi46 from the fungus *Chaetomium globosum*, which was highly upregulated at the early stage of interaction with different pathogens cell wall, and then it was downregulated (Liu et al., 2008). It is important to note that *MfChi* gene expression was altered dramatically within 12 h of exposure to cell walls, indicating a rapid physiological response pathway in *Metschnikowia fructicola* AP47.

As expected, the recombinant chitinase expressed in *P. pastoris* was directly secreted into the medium, with a size of 40.9 kDa, it confirmed to have a high endochitinase activity, and it was the most abundant protein in the medium. These results confirmed MfChi characteristics, previously calculated by sequence analysis tools, and proved that *P. pastoris* is a successful system for yeast protein expression.

The expressed chitinase was able to cause swelling of the hyphae of *Monilinia fructicola* under optical
microscope, confirming the reliability of dual culture procedure to evaluate the presence of active
hydrolases *in vitro*.

Previous studies confirmed that chitinase can decompose fungal cell walls (Li et al., 2005; Liu et al., 2008) since the chitin is the essential cell wall component of many fungal pathogens (Liu et al., 2012b). This study demonstrated that the recombinant chitinase MfChi expressed in *P. pastoris* is highly effective in reducing spore germination and germ tube length of *Monilinia* spp. *in vitro*, but its antifungal activity mainly depends on the chitinase concentration.

In vivo trials confirmed the high efficacy shown in vitro by the recombinant chitinase, and 450 451 demonstrated the capacity of the chitinase to keep its activity for some days in the unfavourable environment of the fruit wounds and on fruit stored at room temperature. However, the efficacy was 452 dependent on the enzyme concentration and the temporal distance from the chitinase treatment which 453 454 could be explained by the loss of chitinolytic activity with increasing the number of storage days. Our results are in accordance with the results obtained by previous studies, which demonstrated that the 455 efficacy of recombinant enzymes expressed in *P. pastoris* against pathogens in fruits is dependent on 456 457 the concentration of the enzyme and the time between enzyme treatment and pathogen inoculation (Banani et al., 2014; Yan et al., 2008). 458

Interestingly, though *M. fructicola* has bigger conidial dimension, more abundant sporulation, longer germ tube length and higher growth rates than *M. laxa* (EPPO, 2009), the recombinant chitinase similarly controlled both species, either *in vitro* or *in vivo*, showing that its efficacy is not dependent on the pathogen species.

To our knowledge, it is the first time that a chitinase gene from the yeast *Metschnikowia fructicola* was cloned and characterized. The present work provides the clarification of the chitinase role in the antagonistic activity of the biocontrol agent *Metschnikowia fructicola* AP47.

Moreover, our results confirm that MfChi chitinase has an excellent antifungal activity to control *Monilinia* species, present as postharvest pathogens not only on stone fruits, but also onother fruits such as apples and pears. *Pichia pastoris* KM71 is a suitable strain for the expression of foreign chitinase genes, which could facilitate the development of a new cost-effective technique for largescale production of recombinant chitinases for biocontrol of fungal postharvest pathogens of fruit.

This work shows that the chitinase MfChi could be developed as a postharvest treatment with antimicrobial activity for fruit undergoing a short shelf life, since it is able to keep its enzymatic activity for some days on the fruit surface and in the wounds. The storage conditions tested in the experiments are highly favourable to the development of brown rot, while when peaches are stored at cold storage temperature, the disease development is slower and the efficacy of chitinase could be for longer periods. Further work will aim at determining the best conditions of activity and stability of this enzyme to obtain the maximum efficacy against the pathogens.

478

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483

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# 645 Supplementary Tables

Parameters**	MfChi 365	
Number of amino acids		
Molecular weight (kDa)		40.92
Theoretical pI		5.27
Amino acids and composition*	Nr	Ср
Ala (A)	25	6.8%
Arg (R)	16	4.4%
Asn (N)	29	7.9%
Asp (D)	34	9.3%
Cys (C)	7	1.9%
Gln (Q)	15	4.1%
Glu (E)	13	3.6%
Gly (G)	27	7.4%
His (H)	4	1.1%
Ile (I)	10	2.7%
Leu (L)	31	8.5%
Lys (K)	24	6.6%
Met (M)	12	3.3%
Phe (F)	23	6.3%
Pro (P)	17	4.7%
Ser (S)	21	5.8%
Thr (T)	16	4.4%
Trp (W)	3	0.8%
Tyr (Y)	14	3.8%
Val (V)	24	6.6%

# 646 **Supplementary Table 1** Amino acid characterization of *MfChi* of *Metschnikowia fructicola* AP47.

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649 \*\*The parameters of MpChi and MfChi were charachterized with ExPASy Proteomics Server at the

650 website: <u>http://www.expasy.org</u>

<sup>648 \*</sup> Nr: number of residues; Cp: composition (percentage) of each amino residue.

# **Supplementary Table 2** Full name and accession number of the chitinases used for the

653 phylogenetic tree analysis of MfChi, retrieved from NCBI and UniProt database.

Abbreviation	Full names of the chitinases
<i>B. bassiana</i> endo-Chi	Beauveria bassiana, endochitinase gb AAN41260.1
B. circulans ChiA	Bacillus circulans chitinase A1 (chiA) gb M57601.1 BACCHIA3
B. licheniformis Chi	Bacillus licheniformis, chitinase gene, gb   AY205293.1
<i>B. subtilis</i> Chi	Bacillus subtilis chitinase (chi) gene, gb   AF069131.1
C. albicans Chi	Candida albicans, chitinase gb   AAS66201.1
C. albicans  S65110  Chi	Candida albicans, chitinase (EC 3.2.1.14) gb S65110
C. albicans SC5314 Chi	Candida albicans SC5314, chitinase ref XP_719348.1
C. albicans SC5314 Cht2	Candida albicans SC5314, chitinase Cht2 ref XP_721807.1
C. albicans SC5314 Cht2(2)	Candida albicans SC5314, chitinase Cht2 ref XP_721966.1
<i>E. americana</i> Chi	Ewingella americana, chitinase emb X90562.1
I. farinosa Chi	Isaria farinosa, chitinase gb ABD64606.1
<i>I. fumosorosea</i> endo-Chi	Isaria fumosorosea, bacterial-type endochitinase gb AAX19146.1
L. lecanii acidic-Chi	Lecanicillium lecanii, acidic chitinase gb   AAX56960.1
L. lecanii basic-Chi	Lecanicillium lecanii, basic chitinase gb AAV98691.1
<i>M. anisopliae</i> Chi	Metarhizium anisopliae, chitinase gb   AAY32603.1
<i>M. flavoviride</i> Chi	Metarhizium flavoviride, chitinase emb CAB44709.1
Malus x domestica class II CHTMA	Malus x domestica, class II chitinase (CHTMA) gb HQ416905.1
<i>N. rileyi</i> Chi	Nomuraea rileyi , chitinase  AAP04616.1
<i>N. tabacum</i> endo-Chi	Nicotiana tabacum, Acidic endochitinase sp P17514 CHIQ_TOBAC Q
O. sativa CHI11	<i>Oryza sativa</i> subsp. <i>japonica</i> , Chitinase 11 sp Q10S66 CHI11_ORYSJ
S. cerevisiae endo-Chi 2	Saccharomyces cerevisiae endochitinase gb   AAA34539.1
S. cerevisiae endo-Chi1	Saccharomyces cerevisiae endochitinase gb AAA34538.1
S. cerevisiae S288c Cts1p	Saccharomyces cerevisiae S288c, Cts1p ref NP_013388.1
S. cerevisiae RM11-1a endo-Chi	Saccharomyces cerevisiae RM11-1a, endochitinase gb EDV08610.1
S. cerevisiae YJM789 endo-Chi	Saccharomyces cerevisiae YJM789, endochitinase gb   EDN59372.1
S. Suprus Chi	
Streptomyces sp. CNIN	Streptomyces sp. ABKIINW 18 CNIN gene, gb HM/48586.1
<i>I. aureoviride</i> endo-Chi	Trichoderma aureoviride, 42 kDa endochitinase gb AY850032.1
<i>U. dioica</i> endo-Chi	Urtica dioica, Lectin/endochitinase 1 sp P11218  AGI_URTDI

#### 656 **Figure captions**

**Fig. 1** Relative expression levels (transcript accumulation) determined by Reverse transcriptionquantitative real-time polymerase chain reaction (RT-qPCR) of the chitinase gene *MfChi* in *Metschnikowia fructicola* AP47 cultured with *Monilinia fructicola* CWP and without CWP (control) at each time point (6 h, 12 h, 24 h and 48 h). Values were normalized to the control at 6h arbitrarily set to unity. Vertical lines represent the standard error for an average of three biological replicates. Different letters above the columns indicated a significant difference determined by Duncan's Multiple comparison Test (p < 0.05).

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Fig. 2 SDS-PAGE analysis of the recombinant chitinase expressed in *P. pastoris*. Supernatants of the yeast culture were taken from different isolates after 120 h of induction. Lanes: M: molecular weight marker (Precision Plus Protein Dual Color Standards, BIO RAD); 0: *P. pastoris* KM71 isolate transformed with pPIC9 (Control); 1, 2, 3 and 4: some transformed *P. pastoris* isolates with pPIC9-*MfChi*.

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**Fig. 3** Effect of the recombinant chitinase MfChi on spore germination (**A**) and germ tube length (**B**) of *M. fructicola* and *M. laxa* after 9 h and 18 h of incubation at 25 °C in potato dextrose broth medium. The chitinase was applied at 7 ng/ $\mu$ L (C1) and 70 ng/ $\mu$ L (C2). Treatments followed by different letters are statistically different following the Duncan's multiple range test (p < 0.05).

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**Fig. 4** Antifungal activity of the recombinant chitinase MfChi in controlling the decay development of *M. fructicola* and *M. laxa* in wound-inoculated peaches. The chitinase was applied at 7 ng/ $\mu$ L (C1) and 70 ng/ $\mu$ L (C2). The results are the mean of three independent experiments. Treatments followed by different letters are statistically different following the Duncan's multiple range test (p < 0.05).

**Supplementary Fig. 1** Alignment of the deduced amino acid sequence of eight chitinase genes from 681 682 yeasts. The data were retrieved from NCBI database: Candida albicans chitinase (S65110); Candida albicans chitinase (AAS66201.1); Candida albicans SC5314 chitinase (XP\_719348.1); Candida 683 albicans SC5314 chitinase Cht2 (XP\_721807.1); Candida albicans SC5314 chitinase Cht2 684 (XP\_721966.1); Candida tropicalis MYA-3404 chitinase 1 precursor (XP\_002546283.1); 685 Saccharomyces cerevisiae endochitinase (AAA34538.1); Scheffersomyces stipitis CBS 6054 686 687 chitinase (XP 001386607.2). Multiple sequence alignment of proteins was performed by using DNAMAN 7.0. Identical residues are highlighted in black background. The conserved region 688 (LLSLGG and QFYNNYC) marked with asterisks were used to design the degenerate primers to 689 690 amplify the partial sequence of chitinase genes from the antagonistic yeasts Metschnikowia fructicola strain AP47. 691

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693 Supplementary Fig. 2 Nucleotide and deduced amino acid sequences of MfChi. The entire DNA sequence of MfChi is shown together with the corresponding amino acid sequence displayed below 694 695 it. Nucleotides and amino acids are numbered on the left side of the sequence. The start codon (ATG) 696 is underlined with a single line; the stop codon (TGA) is marked with an asterisk; the signal peptide is highlighted with the arrows; four putative *N*-glycosylation sites (NFSN, NLTN, NLTV and NLTN) 697 698 are underlined with double lines, the chitinase catalytic activity site (DGYDFNME) is bolded and underlined with a single line, and the two repeated regions in the 3 prime terminal of the open reading 699 frame (ORF) of *MfChi* are shadowed with grey colour. 700

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Supplementary Fig. 3 Alignment of *MfChi* amino acids with those of chitinase genes from the yeasts,
 *Metschnikowia pulcherrima, Candida albicans, Saccharomyces cerevisiae* and *Scheffersomyces stipitis* with DNAMAN 7.0 and CLASTALW. Chitinase genes for alignments were retrieved from
 NCBI database. Amino acids that are identical between MfChi and other sequences are shadowed
 with black color. Non-coding amino acids were shown in dashed line.

**Supplementary Fig. 4** Alignment of the N-terminal regions of MfChi with those of other known yeast chitinases retrieved from NCBI and UniProt databases. Alignment was performed with DNAMAN 7.0 and CLASTALW. Identical residues are shadowed with black color. Non-coding amino acids are shown in dashed line. Numbers mean the position of selected peptide fragments starting from their corresponding start codons. Six cysteine residues highly conserved are marked with an asterisk. The chitinase family 18 active site is highlighted with a box. The proposed aspartic and glutamic catalytic residues are highlighted with a full-black triangle.

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**Supplementary Fig. 5** Phylogenetic analysis of MfChi and other chitinases from different microorganisms and plants. The amino acid sequences of other chitinases were retrieved from NCBI and UniProt database as seen in Supplementary Table 2. The phylogenetic tree of MfChi was generated using MEGA6 by neighbour-joining method. The numbers at node indicate the bootstrap percentages of 1000 resamples.