

The Nodulation Factor Hydrolase of *Medicago truncatula*: Characterization of an Enzyme Specifically Cleaving Rhizobial Nodulation Signals^{1[W][OPEN]}

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Nodule formation induced by nitrogen-fixing rhizobia depends on bacterial nodulation factors (NFs), modified chitin oligosaccharides with a fatty acid moiety. Certain NFs can be cleaved and inactivated by plant chitinases. However, the most abundant NF of *Sinorhizobium meliloti*, an *O*-acetylated and sulfated tetramer, is resistant to hydrolysis by all plant chitinases tested so far. Nevertheless, this NF is rapidly degraded in the host rhizosphere. Here, we identify and characterize MtNFH1 (for *Medicago truncatula* Nod factor hydrolase 1), a legume enzyme structurally related to defense-related class V chitinases (glycoside hydrolase family 18). MtNFH1 lacks chitinase activity but efficiently hydrolyzes all tested NFs of *S. meliloti*. The enzyme shows a high cleavage preference, releasing exclusively lipodisaccharides from NFs. Substrate specificity and kinetic properties of MtNFH1 were compared with those of class V chitinases from *Arabidopsis thaliana* and tobacco (*Nicotiana tabacum*), which cannot hydrolyze tetrameric NFs of *S. meliloti*. The Michaelis-Menten constants of MtNFH1 for NFs are in the micromolar concentration range, whereas nonmodified chitin oligosaccharides represent neither substrates nor inhibitors for MtNFH1. The three-dimensional structure of MtNFH1 was modeled on the basis of the known structure of class V chitinases. Docking simulation of NFs to MtNFH1 predicted a distinct binding cleft for the fatty acid moiety, which is absent in the class V chitinases. Point mutation analysis confirmed the modeled NF-MtNFH1 interaction. Silencing of *MtNFH1* by RNA interference resulted in reduced NF degradation in the rhizosphere of *M. truncatula*. In conclusion, we have found a novel legume hydrolase that specifically inactivates NFs.

Nitrogen-fixing bacteria (rhizobia) establish a root nodule symbiosis with legume plants. Rhizobial infection and subsequent nodule formation depend on

bacterial signaling, namely nodulation factors (NFs), which are perceived by specific LysM receptor kinases of the host plant (Antolín-Llovera et al., 2012). NFs are lipochitooligosaccharides (i.e. modified chitin oligosaccharides with a fatty acyl moiety replacing the *N*-acetyl group at the nonreducing end). In most cases, the oligosaccharide moiety of NFs consists of four or five β -1,4-linked GlcNAc residues. Depending on the rhizobial strain, NFs possess diverse additional substitutions on the oligosaccharide backbone (Perret et al., 2000). Such structural modifications of NFs influence the binding of NFs to the LysM receptor kinases. Hence, NF signaling in host plants culminating in the subsequent expression of symbiosis-specific genes and nodule formation is only triggered by specific NFs (Perret et al., 2000; Antolín-Llovera et al., 2012).

Due to structural similarities to chitin, various NFs can be hydrolyzed by plant chitinases (Staehelin et al., 1994a, 1994b; Goormachtig et al., 1998; Minic et al., 1998; Schultze et al., 1998; Ovtsyna et al., 2000). Chitinases (EC 3.2.1.14) are defined as enzymes that cleave β -1,4 glycosidic linkages in chitin (poly-GlcNAc) and chitin oligosaccharides (Collinge et al., 1993). Chitin is a major polysaccharide in fungal cell walls and exoskeletons of arthropods. Plants do not possess chitin but use chitinases as defense proteins against chitin-containing antagonists, such as fungi and arthropods.

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Various plant chitinases possess additional lysozyme activity (i.e. they can cleave β -1,4-linkages between *N*-acetylmuramic acid and GlcNAc in bacterial peptidoglycan; Collinge et al., 1993). Furthermore, certain plant chitinases appear to hydrolyze endogenous substrates such as GlcNAc-containing arabinogalactan proteins (De Jong et al., 1992; van Hengel et al., 2001). Based on their amino acid sequence, chitinolytic enzymes of plants belong either to glycoside hydrolase family 18 (class III and V chitinases) or family 19 (class I, II, and IV chitinases; Henrissat, 1991).

Class V chitinases of plants, also classified as pathogen-related protein family 11 (PR-11) proteins, form a separate clade within the phylogenetic tree of family 18 glycoside hydrolases (Karlsson and Stenlid, 2009). A phylogenetic tree of class V chitinases and closely related proteins is shown Supplemental Figure S1. In contrast to other plant chitinase classes, only a few class V proteins have been characterized. Nearly two decades ago, the first class V chitinase (NtChiV; also named Pz chitinase) was purified from tobacco mosaic virus-inoculated leaves of tobacco (*Nicotiana tabacum*). This defense-related enzyme efficiently cleaves chitin oligosaccharides (Heitz et al., 1994; Melchers et al., 1994; Brunner et al., 1998). Recently, enzymatic characterization and protein crystal structure analysis were performed with the NtChiV protein expressed in *Escherichia coli* as well as with AtChiC, a related class V chitinase from *Arabidopsis thaliana* (Ohnuma et al., 2011a, 2011b). Furthermore, a class V chitinase purified from the gymnosperm *Cycas revoluta* exhibits transglycosylation activity at high concentrations of chitin oligosaccharides (Taira et al., 2009, 2010). In addition to these enzymes, a class V chitinase-related lectin (RobpsCRA) from the legume tree *Robinia pseudoacacia* has been characterized. This protein binds to high-Man *N*-glycans but lacks catalytic activity (Van Damme et al., 2007). Similarly, the extracellular domain of the chitinase-related receptor-like kinase 1 of tobacco is related to class V chitinases (Kim et al., 2000).

In nodule symbiosis, host chitinases have the potential to inactivate NFs. When tested in root hair deformation bioassays, the purified cleavage products are at least 1,000-fold less active (Heidstra et al., 1994; Staehelin et al., 1994b). In fact, NFs produced by compatible rhizobia were found to be rapidly degraded in the rhizosphere of legumes (Heidstra et al., 1994; Staehelin et al., 1994b, 1995; Ovtsyna et al., 2000, 2005). NFs with a shorter oligosaccharide backbone and carrying additional chemical substitutions are usually more resistant against degradation by chitinases (Staehelin et al., 1994a, 1994b; Minic et al., 1998; Schultze et al., 1998; Ovtsyna et al., 2000). NodSm-IV(C16:2, Ac, S), an *O*-acetylated and sulfated tetrasaccharide with a C16:2 fatty acid chain (Lerouge et al., 1990), is the most abundant NF produced by *Sinorhizobium meliloti* (Supplemental Fig. S2). This NF is resistant to hydrolysis by all plant chitinases tested so far (Schultze et al., 1998). Nevertheless, when added to roots of the host plant alfalfa (*Medicago sativa*), NodSm-IV(C16:2,

Ac, S) was rapidly degraded to a lipodisaccharide, and increased NF hydrolysis was observed when roots were pretreated with NFs (Staehelin et al., 1995). NFs were also found to induce an NF-cleaving activity in the rhizosphere of pea (*Pisum sativum*) plants, and this induction did not happen in NF signaling-deficient pea mutants (Ovtsyna et al., 2000, 2005). Similarly, chitinolytic activities of certain chitinase isoforms of legumes are induced during symbiosis with rhizobia or in response to purified NFs (Staehelin et al., 1992; Xie et al., 1999; Ovtsyna et al., 2005). Furthermore, transcript levels of the class III chitinase gene *Srchi13* in the tropical legume *Sesbania rostrata* are strongly stimulated during symbiosis, and the encoded protein expressed in *E. coli* can degrade NFs of *Azorhizobium caulinodans* to nonidentified cleavage products (Goormachtig et al., 1998). In the model legume *Medicago truncatula*, expression of the class V chitinase gene *MtChit5* is induced during symbiosis with *S. meliloti* (Salzer et al., 2004). Taken together, these findings raise the question of whether legumes possess chitinase-related enzymes that specifically cleave NFs.

In this study, we show that the MtChit5 protein of *M. truncatula* lacks chitinase activity but efficiently hydrolyzes NFs of *S. meliloti*. Therefore, we renamed MtChit5 to MtNFH1 (for *M. truncatula* Nod factor hydrolase1). A substrate-binding model, supported by point mutation analysis, provides a molecular explanation for the identified MtNFH1-NF interaction.

RESULTS

MtNFH1 Encodes an NF-Cleaving Enzyme

In previous work, we identified a symbiosis-related chitinase V gene, *MtChit5*, in *M. truncatula* ecotype R108-1. The expression of this gene is strongly induced in nodule symbiosis (Salzer et al., 2004). What is the catalytic activity of the enzyme encoded by this gene? To answer this question, the corresponding DNA was cloned into vector pET28b in order to express it as a His-tagged recombinant protein in *E. coli* BL21 (DE3). Two related genes of *M. truncatula* (*Mt75352* and *MtCRA*) known to be expressed in *M. truncatula* (Supplemental Table S1; Supplemental Fig. S1) were also cloned and expressed in *E. coli*. The *Mt75352* sequence encodes a putative class V chitinase, whereas *MtCRA* is related to the previously characterized lectin RobpsCRA of *R. pseudoacacia* (Van Damme et al., 2007). For comparison, the class V chitinases AtChiC of *Arabidopsis* (Ohnuma et al., 2011a) and NtChiV of tobacco (Ohnuma et al., 2011b) were cloned and expressed in a similar way. AtChiC and NtChiV show significant amino acid sequence homology to MtChit5 (sequence identity, 42% and 39%, respectively). The recombinant proteins were isolated by nickel affinity purification and analyzed by SDS-PAGE. Rabbit antiserum raised against the recombinant MtChit5 (renamed MtNFH1; see below) was cross-reactive with the other four proteins, indicating successful purification of the recombinant proteins (Fig. 1A).

Purified proteins were then used for enzyme assays with pentameric and tetrameric NFs (Supplemental Fig. S2). These substrates had been HPLC purified from *S. meliloti* strain 1021 (pEK327) (Schultze et al., 1992). We found that the purified product of the *MtChit5* gene rapidly hydrolyzed all the NFs (Table I) and therefore renamed it MtNFH1. Representative HPLC results with NF substrates and acylated cleavage products (lipooligosaccharides after *n*-butanol extraction)

are shown in Figure 1B. The lipodisaccharide NodSm-II (C16:2) was formed when either pentameric NodSm-V (C16:2, S) or tetrameric NodSm-IV(C16:2, S) was used as a substrate (Fig. 1B, a and b). NodSm-IV(C16:2), a chemically desulfated NF, was also hydrolyzed by MtNFH1, albeit to a lesser extent (Fig. 1B, c). Interestingly, MtNFH1 was also able to cleave the *O*-acetylated tetramer NodSm-IV(C16:2, Ac, S); the more slowly migrating *O*-acetylated lipodisaccharide

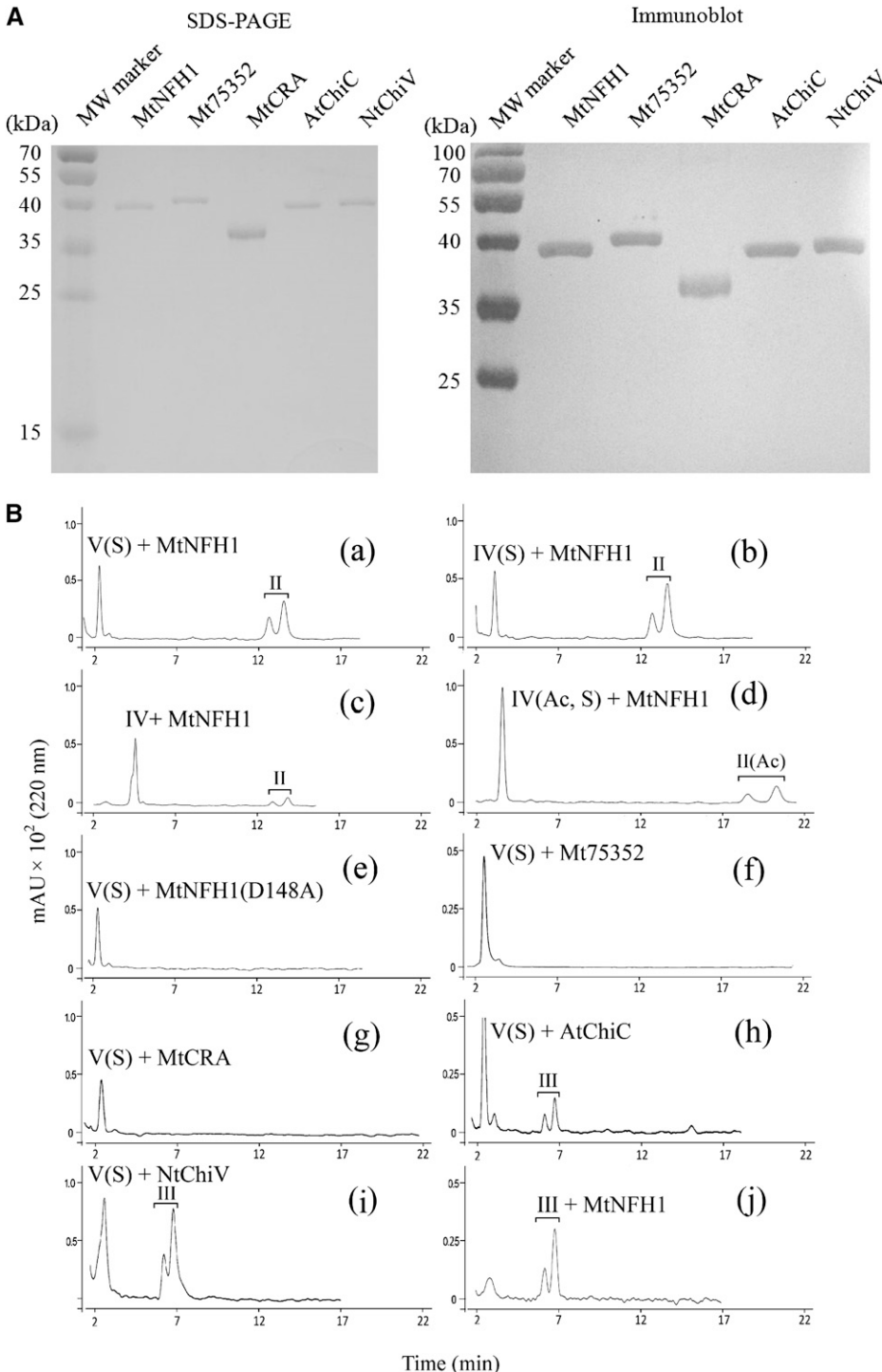


Figure 1. Purification of recombinant proteins and activity test with NF substrates. A, SDS-PAGE analysis (left) and immunoblot detection (right) of His-tagged MtNFH1, Mt75352, MtCRA, AtChiC, and NtChiV expressed in *E. coli* BL21 (DE3). Purified proteins (1 μ g) were analyzed on SDS-PAGE gels stained with Coomassie Brilliant Blue R-250. Immunoblot analysis was performed with a rabbit serum raised against MtNFH1 and the 3,3'-diaminobenzidine reagent. MW, Molecular mass. B, Separation of purified NFs and acylated cleavage products on a Nova Pak C18 column. The NF substrates were incubated with the indicated proteins at 37°C, extracted with *n*-butanol, and subjected to reverse-phase HPLC analysis. The cleavage products were separated into anomers (double peaks). Chromatograms are as follows: MtNFH1 (0.9 μ g mL⁻¹) incubated with 40 μ M NodSm-V(C16:2, S) in 0.1 mL for 20 min (a), with 40 μ M NodSm-IV(C16:2, S) in 0.1 mL for 40 min (b), with 20 μ M desulfated NodSm-IV(C16:2) in 0.1 mL for 120 min (c), with 60 μ M NodSm-IV(C16:2, Ac, S) in 0.1 mL for 40 min (d); the mutant protein MtNFH1(D148A) (0.9 μ g mL⁻¹) incubated with 20 μ M NodSm-V(C16:2, S) in 0.1 mL for 4 h (e); Mt75352 (0.9 μ g mL⁻¹) incubated with 20 μ M NodSm-V(C16:2, S) in 0.1 mL for 4 h (f); MtCRA (3 μ g mL⁻¹) incubated with 20 μ M NodSm-V(C16:2, S) in 0.1 mL for 4 h (g); AtChiC (3 μ g mL⁻¹) incubated with 40 μ M NodSm-V(C16:2, S) in 0.1 mL for 2 h (h); NtChiV (3 μ g mL⁻¹) incubated with 40 μ M NodSm-V(C16:2, S) in 0.1 mL for 2 h (i); and MtNFH1 (0.9 μ g mL⁻¹) incubated with 20 μ M NodSm-III (C16:2, S) in 0.1 mL for 4 h (j); II, NodSm-II(C16:2); II(Ac), NodSm-II(C16:2, Ac); III, NodSm-III(C16:2); IV, NodSm-IV(C16:2); IV(S), NodSm-IV(C16:2, S); IV(Ac, S), NodSm-IV(C16:2, Ac, S); V(S), NodSm-V(C16:2, S).

NodSm-II(C16:2, Ac) was detected by HPLC (Fig. 1B, d). Fractions containing lipodisaccharides were collected, and the chemical structures of lipodisaccharides were verified by mass spectrometry (Supplemental Fig. S3; Supplemental Table S2).

MtNFH1(D148A), a mutant protein in which the conserved Asp residue Asp-148 of the predicted catalytic center was changed to Ala, did not show enzyme activity (Fig. 1B, e). Mt75352 and MtCRA, proteins of *M. truncatula* related to MtNFH1, were unable to hydrolyze the NF substrates (Fig. 1B, f and g). AtChiC of *Arabidopsis* hydrolyzed pentameric NodSm-V(C16:2, S), but the cleavage products were different: the more rapidly migrating lipotrisaccharide NodSm-III(C16:2) was formed (Fig. 1B, h). Similarly, the tobacco enzyme NtChiV released NodSm-III(C16:2) from the pentameric substrate (Fig. 1B, i), as reported previously (Schultze et al., 1998). The structure of the product NodSm-III(C16:2) was confirmed by mass spectrometry (Supplemental Fig. S3; Supplemental Table S2). As compared with MtNFH1, hydrolytic activities of AtChiC and NtChiV for the substrate NodSm-V(C16:2, S) were more than 100-fold lower (Table I). Tetrameric NFs were neither cleaved by AtChiC nor by NtChiV (Schultze et al., 1998). MtNFH1 was unable to hydrolyze purified NodSm-III(C16:2), providing evidence that the enzyme directly releases lipodisaccharides from the NF substrates (Fig. 1B, j).

MtNFH1 Specifically Cleaves NFs

We further tested whether MtNFH1 can hydrolyze nonmodified chitin oligosaccharides (oligomers of GlcNAc), which likely represent the natural substrates

for AtChiC and NtChiV. The hexamer (GlcNAc)₆ and the pentamer (GlcNAc)₅ were used in enzyme assays with purified proteins. After incubation, chitin oligosaccharides were separated on an amino column. Surprisingly, MtNFH1 protein samples, which had shown such a high activity with NF substrates, completely lacked activity with these nonmodified chitin oligosaccharides (Table I; Fig. 2). To test a possible inhibitory effect of chitin oligosaccharides on NF degradation, MtNFH1 was incubated with NodSm-V(C16:2, S) and a 500-fold higher concentration of (GlcNAc)₅. NFs and acylated degradation products were then analyzed by HPLC. Formation of NodSm-II(C16:2) by MtNFH1 was not affected by (GlcNAc)₅, as shown in Supplemental Figure S4.

In contrast to MtNFH1, purified AtChiC and NtChiV proteins efficiently hydrolyzed nonmodified chitin oligosaccharides in accordance with previous reports (Brunner et al., 1998; Ohnuma et al., 2011a, 2011b). (GlcNAc)₆ was cleaved into (GlcNAc)₃ or into (GlcNAc)₄ and (GlcNAc)₂. The product (GlcNAc)₄ was then further hydrolyzed into (GlcNAc)₂. The substrate (GlcNAc)₅ was cleaved into (GlcNAc)₃ and (GlcNAc)₂ (Table I; Fig. 2).

When colloidal chitin, glycolchitin, and carboxymethylchitin-Remazol Brilliant Violet 5R (CM-chitin-RBV) were tested as substrates, AtChiC and NtChiV clearly showed enzymatic activity, while MtNFH1 was completely inactive (Table I). The Mt75352 and MtCRA proteins were similarly inactive. MtNFH1, like AtChiC and NtChiV, lacked lysozyme activity as measured with *Micrococcus lysodeikticus* cells (Table I), which is in agreement with a previous report on NtChiV (Heitz et al., 1994). NtChiV inhibited hyphal growth of the fungus *Trichoderma viride* (Melchers et al., 1994; Ohnuma et al., 2012), and this was confirmed in a similar bioassay

Table I. Activities of MtNFH1, AtChiC, and NtChiV with the indicated substrates

Substrate	Enzyme Activity ^a			Assay
	MtNFH1	AtChiC	NtChiV	
NodSm-V(C16:2, S)	156.5 ± 8.1	0.37 ± 0.06	1.27 ± 0.23	Reverse-phase HPLC analysis (Nova Pak C18 column) ^b
NodSm-IV(C16:2, S)	121.8 ± 21.7	ND ^c	ND	
NodSm-IV(C16:2, Ac, S)	46.5 ± 4.2	ND	ND	
NodSm-III(C16:2)	ND	ND	ND	Reverse-phase HPLC analysis (amino column) ^d
(GlcNAc) ₆	ND	393.1 ± 26.8	205.9 ± 29.8	
(GlcNAc) ₅	ND	266.0 ± 19.9	171.1 ± 19.0	
Colloidal chitin	ND	0.020 ± 0.004	0.012 ± 0.001	
Glycolchitin	ND	13.49 ± 3.32	7.66 ± 0.22	
CM-chitin-RBV	ND	0.83 ± 0.22 ^e	1.33 ± 0.03	Colorimetric assay
<i>M. lysodeikticus</i> cells	ND	ND	ND	Lysozyme assay

^aData are shown for the indicated 6×His-tagged recombinant proteins. Proteolytic removal of the tag with factor Xa did not affect the enzyme activity of MtNFH1 as determined with NodSm-IV(C16:2, S). Mt75352 and MtCRA lacked enzymatic activity when tested either with NFs or chitinous substrates. Activities were determined at 37°C with a substrate concentration of 150 μM for NFs from *S. meliloti* or the lipotrisaccharide NodSm-III(C16:2), 3.6 mM for (GlcNAc)₆, 4.5 mM for (GlcNAc)₅, approximately 10 mg mL⁻¹ for colloidal chitin, 20 mg mL⁻¹ for glycolchitin, 0.9 mg mL⁻¹ for CM-chitin-RBV, and 0.45 mg mL⁻¹ for *M. lysodeikticus* cells. Data indicate means ± SD from at least three independently purified enzyme preparations. ^bFormation of the lipodisaccharide NodSm-II(C16:2) or NodSm-III(C16:2, Ac) by MtNFH1 and formation of the lipotrisaccharide NodSm-III(C16:2) by AtChiC or NtChiV. ^cND, Not detected. ^dAtChiC and NtChiV degraded (GlcNAc)₆ into (GlcNAc)₃ or (GlcNAc)₂ and (GlcNAc)₄; (GlcNAc)₄ was further degraded into (GlcNAc)₂; (GlcNAc)₅ was cleaved into (GlcNAc)₃ and (GlcNAc)₂. ^eEnzyme activity expressed as ΔA₅₅₀ mg⁻¹ protein s⁻¹.

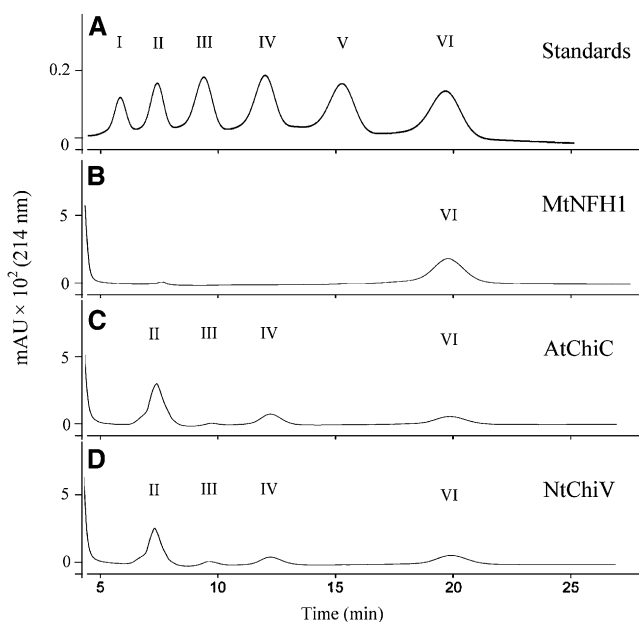


Figure 2. MtNFH1 does not hydrolyze nonmodified chitin oligosaccharides. Examples of HPLC chromatograms showing separation of the substrate (GlcNAc)₆ and cleavage products are presented. After incubation (37°C) with the indicated proteins, reaction mixtures (in 25 mM sodium acetate buffer, pH 5.0; 30- μ L test volume) were diluted with an equal volume of 50% acetonitrile and directly loaded onto the amino column. A, (GlcNAc)_n standards ($n = 1-6$; 10 nmol). B, (GlcNAc)₆ (5.8 mM) incubated with MtNFH1 (4.5 μ g mL⁻¹) for 3 h. C, (GlcNAc)₆ (5.8 mM) incubated with AtChiC (5.0 μ g mL⁻¹) for 30 min. D, (GlcNAc)₆ (5.8 mM) incubated with NtChiV (5.0 μ g mL⁻¹) for 30 min. I, GlcNAc; II, (GlcNAc)₂; III, (GlcNAc)₃; IV, (GlcNAc)₄; V, (GlcNAc)₅; VI, (GlcNAc)₆; mAU, milliabsorbance units.

in this study. The MtNFH1 protein lacked such antifungal activity, however (Fig. 3).

Kinetic Studies Indicate Efficient NF Hydrolysis at Low Substrate Concentrations

Kinetic studies with varying concentrations of NFs were performed to determine K_m and catalytic rate constant (k_{cat}) values of the purified proteins (Table II; Supplemental Fig. S5). MtNFH1 showed K_m values at micromolar concentrations with all tested NFs of *S. meliloti*. A slightly higher K_m and a lower k_{cat} value were determined for the *O*-acetylated substrate NodSm-IV(C16:2, Ac, S) as compared with NFs without an *O*-acetyl group. Similarly, low K_m values were also obtained for AtChiC and NtChiV releasing NodSm-III(C16:2) from NodSm-V(C16:2, S). As compared with MtNFH1, however, only low k_{cat} values were determined.

The results of kinetic studies with AtChiC and NtChiV for the substrates (GlcNAc)₆ and (GlcNAc)₅ are summarized in Supplemental Table S3. Relatively high K_m values in the millimolar range were determined for these substrates. On the other hand, the k_{cat}

values for (GlcNAc)₆ and (GlcNAc)₅ were found to be more than 2 orders of magnitude higher as compared with those for NodSm-V(C16:2, S) shown in Table II.

Docking Simulation Predicts a Specific Binding Cleft for a Fatty Acid Chain

Since MtNFH1 cleaves NFs with their fatty acyl decorations, but not the simple chitin oligosaccharides, we wondered whether the fatty acid chain was required for correct substrate binding. To analyze substrate recognition on a molecular level, a three-dimensional model of MtNFH1 was constructed by homology modeling, using the crystal structures of AtChiC or NtChiV as structural templates. The structures of the substrates NodSm-V(C16:2, S) and (GlcNAc)₅ were then manually docked to the active site of the proteins. Aligned primary sequences of the three proteins and the modeled substrate-protein complexes are shown in Figure 4. All three enzymes possess a predicted substrate-binding pocket for GlcNAc residues. Noteworthy, two loops in MtNFH1 (loops A and B) are predicted to contribute to the formation of the binding cleft for the C16:2 fatty acid moiety. Loop A consists of amino acid residues between strand 9 and helix 6 (GSGS motif), and loop B is formed by residues between strands 12 and 13 (GP GPGVDGG motif; Fig. 4A). In contrast to the AtChiC and NtChiV sequences, these two loops of MtNFH1 are shorter and are Gly rich, which results in a prominent cleft that can accommodate the C16:2 fatty acid moiety of NodSm-V(C16:2, S) (Fig. 4B). Based on our model, NF substrates perfectly fit to MtNFH1 in this way and are then cleaved into NodSm-II(C16:2) and a sulfated chitin oligosaccharide. Hence, MtNFH1 targets NFs at their nonreducing

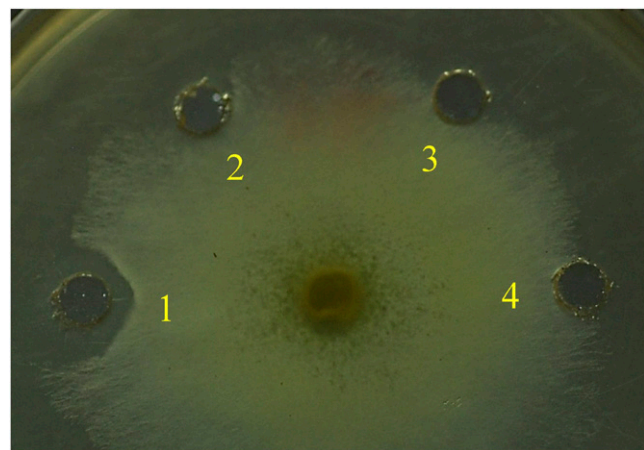


Figure 3. MtNFH1 does not inhibit the growth of the fungus *T. viride*. Purified proteins (3.2 μ g in 50 μ L) of NtChiV (1), AtChiC (2), or MtNFH1 (3) were pipetted into peripheral wells close to a growing mycelium of *T. viride* (inoculation at the center of the agar plate). The sodium acetate buffer (25 mM, pH 5.0) without enzyme was added as a control (4). The plate shown (demonstrating growth inhibition activity for NtChiV) was photographed after an incubation for 14 h at 27°C.

Table II. K_m and k_{cat} values for MtNFH1, AtChiC, and NtChiV with NFs as substrates

Substrate	MtNFH1			AtChiC			NtChiV		
	K_m μM	k_{cat} s^{-1}	$k_{cat} K_m^{-1}$ $mM^{-1} s^{-1}$	K_m μM	k_{cat} s^{-1}	$k_{cat} K_m^{-1}$ $mM^{-1} s^{-1}$	K_m μM	k_{cat} s^{-1}	$k_{cat} K_m^{-1}$ $mM^{-1} s^{-1}$
NodSm-V(C16:2, S)	52.7 ± 2.3^a	8.4 ± 0.6	159.3 ± 17.6	80 ± 6.3	0.033 ± 0.001	0.41 ± 0.02	68 ± 6.4	0.088 ± 0.004	1.3 ± 0.2
NodSm-IV(C16:2, S)	60.1 ± 4.7	7.9 ± 0.2	132.5 ± 13.4	ND ^b	ND	ND	ND	ND	ND
NodSm-IV(C16:2, Ac, S)	97.3 ± 12.2	3.5 ± 0.1	36.4 ± 5.5	ND	ND	ND	ND	ND	ND

^aData indicate means \pm SD from enzyme tests (37°C) with three independently purified protein preparations. ^bND, No activity detected.

ends. On the other hand, there is apparently no space in AtChiC and NtChiV to accommodate the fatty acid moiety at this position (Fig. 4B), which corresponds to the observation that they cannot release NodSm-II (C16:2) from NodSm-V(C16:2, S).

A superimposed model of MtNFH1 and NtChiV indicates changes in sequence/structure that lead to the formation of the fatty acid-binding cleft (Fig. 4C). Specifically, the Tyr and Ser residues of NtChiV (Tyr-236 and Ser-265), and the corresponding residues in AtChiC are substituted in MtNFH1 by a Lys and Gly at the corresponding positions (Lys-241 and Gly-267). These two residues are located at the entrance of the cleft between loops A and B, and these sterically more bulky Tyr and Ser residues in AtChiC and NtChiV are predicted to block the formation of a cleft. Furthermore, NtChiV and AtChiC have an extra Pro residue (Pro-188 in NtChiV) that is absent in loop A of MtNFH1. The deletion of this Pro residue in MtNFH1 results in a shorter loop A and thus enlarges the predicted fatty acid-binding cleft.

Point Mutation Analysis Supports the Modeled NF-MtNFH1 Interaction

Our model predicts that Lys-241 and Gly-267 of MtNFH1, which guard the entrance of the fatty acid-binding cleft, should be important for substrate recognition. To test this hypothesis, we constructed a double mutant of MtNFH1 (K241Y/G267S) in which Lys-241 is substituted by Tyr and Gly-267 is substituted by Ser (corresponding to Tyr-236 and Ser-265 in NtChiV, respectively). When assayed with NFs as substrates, the MtNFH1(K241Y/G267S) protein showed approximately 2.5-fold increases in K_m and approximately 2-fold decreases in k_{cat} which resulted in an approximately 5-fold reduced activity (k_{cat}/K_m ; Table III). These data provide experimental support for the modeled NF-MtNFH1 interaction.

Silencing of MtNFH1 by RNA Interference Increases NF Levels in the Rhizosphere

When adding NFs of *S. meliloti* to roots of alfalfa, they are rapidly degraded by an NF-cleaving extracellular activity (Staehelin et al., 1994b, 1995). To investigate whether MtNFH1 encodes an enzyme that contributes to NF degradation in the rhizosphere, we transformed *M. truncatula* ecotype R108-1 with *Agrobacterium tumefaciens* carrying pCAMBIA-MtNFH1i.

This RNA interference (RNAi) plasmid contains a construct made of the cauliflower mosaic virus (CaMV) 35S promoter in tandem and a fragment of MtNFH1, which is inserted in both the sense and antisense directions. These sense and antisense sequences are separated by a spacer region. Furthermore, the vector carries a GUS gene under the control of the CaMV 35S promoter, which allows it to distinguish between transformed and nontransformed plants. Primary transformants were regenerated from *A. tumefaciens*-infiltrated leaf discs on selective medium containing hygromycin. Young seedlings from these RNAi lines (T2 generation) were first treated for 1 d with 0.1 μM NodSm-IV(C16:2, S) in order to stimulate the NF-cleaving activity in the rhizosphere. The NF degradation test was then performed by incubation in 15 μM NodSm-IV(C16:2, S). After removal of the seedlings, the NFs and the released NodSm-II (C16:2, S) were extracted from the medium by *n*-butanol and quantified by HPLC analysis. Finally, the seedlings were used for the determination of GUS activity, and plants lacking blue staining were excluded from the experiment. Compared with wild-type plants, an up to 5-fold reduction in NF hydrolysis was measured for plants transformed with pCAMBIA-MtNFH1i, indicating that silencing of MtNFH1 increases NF levels in the rhizosphere (Fig. 5).

DISCUSSION

In this work, we have biochemically characterized MtNFH1, which is encoded by a symbiosis-stimulated *M. truncatula* gene (Salzer et al., 2004). The enzyme shows sequence similarities to class V chitinases but lacks chitinase activity. Instead, MtNFH1 cleaves NFs of the microsymbiont *S. meliloti*. Notably, MtNFH1 hydrolyzes NodSm-IV(C16:2, Ac, S), which is structurally rather different from nonmodified chitin oligosaccharides. Based on the capacity to release lipodisaccharides from NFs (this study) and the increase of MtNFH1 transcripts in NF-treated roots (Salzer et al., 2004), it is likely that MtNFH1 is an ortholog of the previously identified NF-cleaving enzyme of alfalfa, which could be partially purified from root exudates due to its concanavalin A-binding properties (Staehelin et al., 1995). It is likely that MtNFH1 is also a secreted N-glycosylated protein. This assumption is supported by the prediction of an N-terminal signal peptide and four potential N-glycosylation sites in MtNFH1 (Supplemental Table S1)

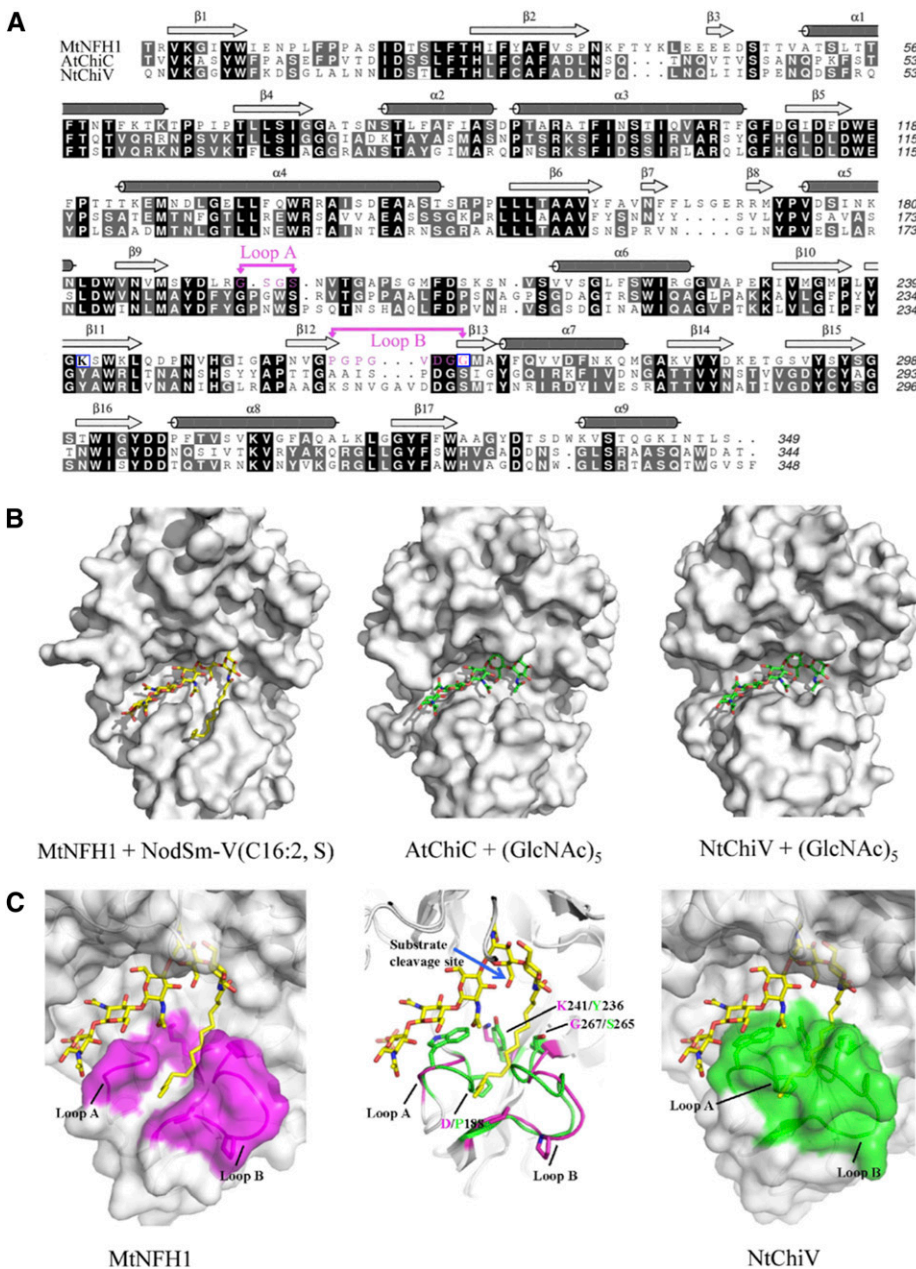


Figure 4. Homology modeling and docking simulation of NodSm-V(C16:2, S) to MtNFH1. **A**, Structure-based sequence alignment of MtNFH1 with AtChiC and NtChiV. Identical amino acid residues are shown on a black background, homologous residues on a gray background, and dashes indicate gaps; α -helices and β -strands are marked above the sequences. Amino acid residues of loops A and B in MtNFH1 are marked in magenta and the point-mutated residues K241 and G267 are framed in blue. **B**, Models of MtNFH1 with NodSm-V(C16:2, S), AtChiC with (GlcNAc)₅, and NtChiV with (GlcNAc)₅. **C**, MtNFH1 (magenta, left) has a prominent cleft between loops A and B that can bind the fatty acid moiety of NodSm-V(C16:2, S). NtChiV (green, right) and AtChiC lack such a binding cleft. The structures of MtNFH1 (magenta) and NtChiV (green) are superimposed to illustrate the expected conformational differences (middle). The cleavage site of NodSm-V(C16:2, S) hydrolyzed by MtNFH1 is indicated by a blue arrow.

as well as our finding that *MtNFH1* RNAi-silenced plants show reduced NF-cleaving activity in the rhizosphere (Fig. 5).

After the initiation of NF signaling, MtNFH1 probably inactivates excess amounts of NFs, as has been suggested for a related hydrolase activity of alfalfa (Stahelin et al., 1994b, 1995), Srchi13 of *S. rostrata* (Goormachtig et al., 1998), and an unknown pea enzyme (Ovtsyna et al., 2000, 2005). Such a feedback response might control the degree of rhizobial invasion into host cells at different symbiotic stages. NFs can trigger responses on root hairs at nanomolar to picomolar concentrations. However, for developing infection threads and for initiating nodules, it has been proposed that higher NF

levels that are present over a longer period of time are required (Perret et al., 2000). As the K_m values of MtNFH1 for NFs are in the micromolar concentration range, we suggest that the substrate concentration for the enzyme is not saturated under natural conditions. Future work is required to examine whether NF levels in the rhizosphere have an impact on establishing symbiosis with *S. meliloti*. Because our RNAi lines show reduced NF-cleaving activity, we currently test them for their effects on nodule formation.

Most plants produce a large set of various chitinases, and this diversity may reflect the importance of these proteins during plant defense against chitinous pathogens, particularly fungi. However, the evolutionary

Table III. Differences of K_m and k_{cat} values of MtNFH1(K241Y/G267S) and wild-type MtNFH1

Protein	NodSm-V(C16:2, S)			NodSm-IV(C16:2, S)		
	K_m	k_{cat}	$k_{cat} K_m^{-1}$	K_m	k_{cat}	$k_{cat} K_m^{-1}$
MtNFH1	100 ^a	100	100	100	100	100
MtNFH1(K241Y/G267S)	271.7 ± 2.3 ^b	54.6 ± 0.7	19.9 ± 0.4	245 ± 1.7	51.6 ± 4.3	21 ± 1.7

^aFor absolute values, see Table II. ^bData (relative values in relation to the wild-type protein; expressed in percentages) indicate means ± SD deduced from enzyme assays (37°C) using three independently purified protein preparations and various amounts of NodSm-V(C16:2, S) and NodSm-IV(C16:2, S), respectively.

driving force for this diversity remains unclear, and it has been proposed that plant chitinases are coevolving with chitinase inhibitors from pathogens (Rausher, 2001). Here, we show that MtNFH1 lacks chitinase activity and instead cleaves NFs of *S. meliloti*. This can be considered as a typical example of neofunctionalization (i.e. adaptation of an enzyme to a novel microbial substrate reminiscent to receptor-ligand coevolution). In fact, class V chitinases are present in ancient land plants, including the primitive gymnosperm *C. revoluta* (Taira et al., 2009, 2010). We suggest that MtNFH1 evolved from an ancestral defense-related chitinase and that gene duplication opened the possibility to develop a symbiotic enzyme that has the capacity to inactivate short and structurally modified NFs such as NodSm-IV(C16:2, Ac, S). In contrast to MtNFH1, recombinant MtCRA and Mt75352 were unable to hydrolyze NFs and may possess unknown nonsymbiotic functions. MtCRA is likely an ortholog of RobpsCRA, which is a nonchitinolytic lectin with binding preference for the pentasaccharide core structure of high-Man *N*-glycans (Van Damme et al., 2007). It remains to be seen whether MtCRA and perhaps MtNFH1 possess similar carbohydrate-binding activities.

NF receptors of legumes possess extracellular LysM domains, which strongly interact with NFs and only weakly with nonmodified chitin oligosaccharides (Broghammer et al., 2012). Similarly, the symbiotic ectoapyrase (lectin nucleotide phosphohydrolase) of the legume *Dolichos biflorus* shows a higher binding affinity for NFs than for chitin oligosaccharides, as determined by a competitive chitin-binding assay (Etzler et al., 1999). In this study, we found that MtNFH1 perfectly discriminates between NFs and chitin oligosaccharides. Accordingly, the K_m values of MtNFH1 for NFs are in the micromolar concentration range, whereas chitin oligosaccharides are neither substrates nor inhibitors. In contrast, the class V chitinases AtChiC and NtChiV can only hydrolyze the pentameric NF, NodSm-V(C16:2, S). In comparison with MtNFH1, their activities degrading this NF are low due to low k_{cat} values. Interestingly, as reflected by the low K_m value for NodSm-V(C16:2, S), AtChiC and NtChiV show significantly higher affinities for this NF than for chitin oligosaccharides (Table II; Supplemental Table S3). These data suggest a further possible role for class V chitinases and related proteins, namely to cleave NF-like signals of

mycorrhizal fungi. Roots of tobacco and *M. truncatula*, but not Arabidopsis, can establish symbiosis with arbuscular mycorrhizal fungi, and a set of fungal lipochitooligosaccharides have been identified recently (Maillet et al., 2011). However, MtNFH1 expression was found to be induced in *S. meliloti*-inoculated *M. truncatula* roots, whereas transcript levels remained low in mycorrhizal roots colonized by the fungus *Glomus intraradices* (Salzer et al., 2004).

Crystallization of AtChiC and NtChiV (Ohnuma et al., 2011a, 2011b) provided useful structural templates for homology modeling of the MtNFH1 protein structure. Docking of NFs to MtNFH1 resulted in a model in which the C16:2 fatty acid moiety fits to a specific binding cleft and apparently helps to bring the carbohydrate moiety in the catalytic pocket to a correct position. The point mutant MtNFH1(D148A) lacked enzyme activity (Fig. 1B, e), indicating that the canonical DXDXE motif of functional glycosyl hydrolase family 18 enzymes (Karlsson and Stenlid, 2009) is also critical for the hydrolysis of NFs. Furthermore, kinetic data of a MtNFH1 mutant protein, modified in the region of the predicted binding cleft for the fatty acid chain (Table III), provide support for the modeled MtNFH1-NF interaction. It is worth noting that the

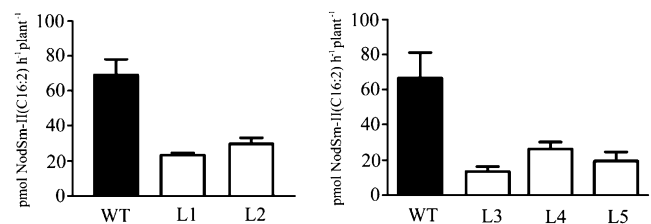


Figure 5. Hydrolysis of NodSm-IV(C16:2, S) by roots of *M. truncatula* seedlings transformed with *A. tumefaciens* carrying pCambia-MtNFH1i. Roots of seedlings (T2 generation) from five independent RNAi lines (named L1–L5) and wild-type plants (WT) were individually pretreated with 0.1 μM NodSm-IV(C16:2, S) for 1 d and then incubated with 15 μM NodSm-IV(C16:2, S) for 18 h. The NF substrate and formed NodSm-II(C16:2) were extracted from the medium with *n*-butanol and separated by reverse-phase HPLC using a Nova Pak C18 column. Data indicate means ± SD from two representative test series (in total, 24 transgenic and seven wild-type plants). NF-cleaving activities of all five RNAi lines are significantly reduced as compared with wild-type plants (Kruskal-Wallis test; $P < 0.03$).

predicted torsion angle between the sugar moiety and the fatty acid chain in the MtNFH1-NF interaction is similar to the corresponding torsion angle in NF-receptor interaction models, in which NFs of *S. meliloti* were individually docked to each of the three LysM domains of a Nod factor perception receptor of *M. truncatula* (Mulder et al., 2006).

It is particularly intriguing that MtNFH1 completely lacks activity when incubated with nonmodified chitin oligosaccharides. As chitin oligosaccharides are smaller than NFs, they likely reach the catalytic center of MtNFH1 without steric hindrance. However, the predicted carbohydrate-binding pocket of MtNFH1 is bigger than that of AtChiC or NtChiV (Fig. 4B), and there is apparently less steric hindrance to fix the flexible chitin oligosaccharide chain into the correct orientation of the active site of MtNFH1. In other words, chitin oligosaccharides cannot adopt a correct conformation required for catalysis. In contrast, NFs with their C16:2 fatty acid moiety fully occupy the substrate-binding pocket in the correct orientation and exactly present the scissile bond to the active site residues of MtNFH1.

MATERIALS AND METHODS

Gene Cloning

DNA sequences encoding proteins without predicted signal peptides from *Medicago truncatula* ecotype R108-1 (MtNFH1, Mt75352, and MtCRA; Supplemental Table S1), *Arabidopsis* (*Arabidopsis thaliana*) ecotype Columbia (AtChiC; accession no. BT029539) and tobacco (*Nicotiana tabacum* 'Xanthi' [NtChiV]; accession no. X78325) were cloned into the expression vector pET28b (Novagen, Merck) and verified by sequencing. MtNFH1 without signal peptide sequence was also cloned into vector pET30 as described by the supplier of the LIC Cloning Kit (Novagen). PCR-based site-directed mutagenesis techniques were used for the introduction of point mutations into MtNFH1. Primers used and plasmids constructed are shown in Supplemental Tables S4 and S5, respectively.

To construct an RNAi plasmid for the silencing of MtNFH1 in *M. truncatula*, a 667-bp fragment of MtNFH1 (MtNFH1 Δ 1-94/ Δ 761-1152) was cloned into pBS-RNAi in the sense and antisense orientations (separated by a 360-bp spacer; Limpens et al., 2004). The primers and plasmid constructs are shown in Supplemental Tables S4 and S5, respectively. Finally, the entire 2,604-bp cassette [containing a double CaMV 35S promoter, the 667-bp fragment in the sense orientation, the spacer, the 667-bp fragment in the antisense orientation and a poly(A) terminator] was cloned into the binary vector pCambia1305.1, which contains a *GUS* gene under the control of the CaMV 35S promoter (<http://www.cambia.org/>). The resulting RNAi plasmid, named pCambia-MtNFH1i, was used for the transformation of *Agrobacterium tumefaciens* strain EHA105 by electroporation.

Phylogenetic Analysis

For the identification of related sequences, the amino acid sequence of MtNFH1 was used as a query sequence for a protein BLAST (BLASTP algorithm) database search (Altschul et al., 1990) at the National Center for Biotechnology Information home page (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>). By selecting the protein database parameter "green plants," a total of 91 related sequences were obtained. The Mt59366, Mt75352, and Mt59600 sequences were extracted from the *M. truncatula* Genome Project database (Mt3.5 CDS database) by using MtNFH1 as a query sequence in a nucleotide Blast (BLASTN algorithm) similarity search (<http://blast.jcvi.org/er-blast/index.cgi?project=mtbe>). The nucleotide sequences were then translated into predicted amino acid sequences using the splicing prediction software SplicePredictor (<http://bioservices.usd.edu/splicepredictor/>). The unrooted phylogenetic tree was then constructed with the MEGA5 program using the

neighbor-joining method (Tamura et al., 2011). The evolutionary distances were computed using the Poisson correction method.

Protein Expression and Purification

Escherichia coli BL21 (DE3) cells carrying the constructed plasmids were cultivated in Luria-Bertani medium, and recombinant protein expression was induced by 0.4 mM isopropyl- β -D-thiogalactopyranoside (18°C for 20 h). Proteins were purified by nickel affinity chromatography with nickel-nitrilotriacetic acid agarose resin beads based on the manufacturer's protocol for protein purification under denaturing or native conditions (Qiagen). Where indicated, the His tag was removed from MtNFH1 [purified from *E. coli* BL21 (DE3) carrying pET30-MtNFH1] by using the protease factor Xa according to the supplier's recommendations (New England Biolabs).

Protein Analysis and Antibodies

Purified proteins were separated by SDS-PAGE, and gels were stained with Coomassie Brilliant Blue R-250. Protein contents of protein bands were then compared with bovine serum albumin standards. In addition, protein contents were photometrically measured according to the method of Bradford (1976). His-tagged MtNFH1 protein, purified under denaturing conditions according to the manufacturer's protocol (Qiagen), was used for immunization of a New Zealand rabbit (provided by the Experimental Animal Centre of Sun Yat-sen University of Medical Science, Guangzhou, China). For immunoblot analysis, proteins were separated by SDS-PAGE and then electrophoretically transferred onto nitrocellulose membranes (Schleicher & Schuell BioScience). Membranes were incubated with the polyclonal rabbit antiserum against MtNFH1 (1:7,000 dilution) and goat anti-rabbit IgG antiserum coupled to horseradish peroxidase. Finally, blots were developed with 3,3'-diamino-benzidine according to the supplier's recommendations (Boster).

Purification of NFs

NodSm-V(C16:2, S), NodSm-IV(C16:2, S), and NodSm-IV(C16:2, Ac, S) were purified from *Sinorhizobium meliloti* 1021 (pEK327) (Schultze et al., 1992). Bacteria were grown in modified glucose-Tris-sodium succinate medium (Kiss et al., 1979), and NFs were purified by HPLC according to previously described procedures (Schultze et al., 1992; Staehelin et al., 1994b, 1995). Briefly, culture supernatants (adjusted to pH 7.5 with HCl) were extracted with *n*-butanol and concentrated under reduced pressure. For the purification of NodSm-IV(C16:2, Ac, S), NFs were fractionated by reverse-phase HPLC (Nova Pak C18, 3.9 \times 150 mm, particle size 4 μ m; Waters) under isocratic conditions using 30% (v/v) acetonitrile/water containing 40 mM ammonium acetate as the mobile phase. NodSm-IV(C16:2, Ac, S), corresponding to a more slowly migrating peak, was collected and the sample dried under reduced pressure. For the purification of NodSm-V(C16:2, S) and NodSm-IV(C16:2, S), the NFs were first completely deacetylated (incubation in 50 mM Tris-HCl, pH 9.0, for 8 h) and then HPLC purified under the same conditions. Finally, NFs were desalted on a Polyosil C18 column (Staehelin et al., 2000). Desulfated NodSm-IV(C16:2) was obtained from NodSm-IV(C16:2, S) by mild acid hydrolysis in methanol-HCl followed by HPLC purification (Demont et al., 1993).

Enzyme Assays with NFs

Purified proteins and NFs at specified amounts were incubated in 25 mM sodium acetate buffer (pH 5.0) at 37°C. Nondegraded NFs and the acylated cleavage products (lipodisaccharides and lipotrisaccharides) were extracted with an equal volume of double-distilled *n*-butanol and dried in a Speed-Vac evaporator. Samples were taken up in 1 μ L of dimethyl sulfoxide (DMSO) and analyzed by reverse-phase HPLC (Nova Pak C18; Waters) using 36% (v/v) acetonitrile/water containing 40 mM ammonium acetate as the mobile phase (Staehelin et al., 1994b). HPLC-purified cleavage products released from NFs were analyzed by mass spectrometry. For measurement of activities and determination of kinetic parameters, the incubation times varied to ensure that the product formation was less than 25% of the substrate amount. Kinetic parameters were deduced by using GraphPad Prism version 5.00 (GraphPad Software).

An NF hydrolysis assay with intact roots was performed as described previously (Staehelin et al., 1994b, 1995). Briefly, experiments were performed with young seedlings from wild-type plants of *M. truncatula* ecotype R108-1

and *MtNFH1* RNAi-silenced lines (five lines, T2 generation, nine to 19 single plants per line). Seedlings were first incubated in 1-mL syringes filled with 400 μ L of Jensen medium (Van Brussel et al., 1982) containing 0.5% (v/v) DMSO and 0.1 μ M NodSm-IV(C16:2, S) at 24°C in the dark. The following day, the pretreated seedlings were transferred to new 1-mL syringes filled with 400 μ L of Jensen medium containing 0.5% (v/v) DMSO and 15 μ M NodSm-IV(C16:2, S). After incubation at 24°C for 18 h in the dark, seedlings were removed and plants from RNAi lines were stained for GUS activity. NodSm-IV(C16:2, S) and formed NodSm-II(C16:2) were extracted from the medium with an equal volume of distilled *n*-butanol. Dried samples were resuspended in 1 μ L of DMSO and subjected to HPLC analysis as described above. Data were statistically analyzed with the nonparametric Kruskal-Wallis test, which is suitable for unequal replications.

Mass Spectrometric Analysis

HPLC fractions containing NodSm-II(C16:2), NodSm-II(C16:2, Ac), or NodSm-III(C16:2) were collected, and samples were dried and desalted on a Polygosil C18 column. Matrix-assisted laser-desorption/ionization (MALDI)-time of flight (TOF) analysis was performed with an Ultraflex III MALDI-TOF/TOF mass spectrometer (Bruker Daltonics) in positive ionization mode. Samples were ionized with a smartbeam I UV laser (at a wavelength of 355 nm). Samples were then resuspended in sterilized water and mixed with the matrix (5 mg mL⁻¹ 2,5-dihydroxybenzoic acid in 30% acetonitrile). The matrix:sample mixture ratio was 1:1, and the final volume was 2 μ L. The mixture was applied to an Anchorchip probe (Bruker Daltonics) and air dried. Observed ion masses corresponded to [M+H]⁺ and [M+K]⁺ due to protonation and the formation of potassium adducts.

Enzyme Assays with Chitin Oligosaccharides

(GlcNAc)₂, (GlcNAc)₃, (GlcNAc)₄, (GlcNAc)₅, and (GlcNAc)₆ were purchased from Seikagaku Kogyo. Enzyme activities were tested by incubating chitin oligosaccharides with different enzymes in 25 mM sodium acetate buffer, pH 5.0, at 37°C. After incubation, reaction mixtures were diluted with an equal volume of 50% acetonitrile and then loaded onto a μ Bondapak NH₂ HPLC column (0.8 \times 10 cm; Waters) using 67% (v/v) acetonitrile/water as the mobile phase. The flow rate was 1 mL min⁻¹, and the oligosaccharides were detected by ultraviolet A₂₁₄. For measurement of enzyme activities and determination of kinetic parameters, the incubation times varied to ensure that the amount of product formation was less than 25% of the substrate amount. Kinetic parameters were deduced with the help of GraphPad Prism version 5.00 (GraphPad Software).

Enzyme Assays with Chitinous Substrates

All assays were performed at 37°C with specified amounts of purified His-tagged proteins in 25 mM sodium acetate buffer (pH 5.0) and substrate concentrations indicated in Table I. Colloidal chitin was obtained by acetylation of chitosan (Sigma-Aldrich) using acetic anhydride (Molano et al., 1977). Recombinant *N*-acetylglucosaminidase from *M. truncatula* was used to convert released oligosaccharides into GlcNAc, which was quantified with a *p*-dimethylaminobenzaldehyde solution (Ehrlich's reagent; Reissig et al., 1955). Glycolchitin was obtained by acetylation of glycol chitosan (Sigma-Aldrich) with acetic anhydride (Molano et al., 1977; Trudel and Asselin, 1990). Reducing end sugars of hydrolyzed glycolchitin were measured with the Lever assay (Lever, 1972). CM-chitin-RBV was bought from Loewe Biochemica. The enzyme activity was photometrically determined based on the manufacturer's instructions (Wirth and Wolf, 1990).

Lysozyme Assay and Fungal Growth Inhibition Test

Lyophilized *Micrococcus lysodeikticus* cells were purchased from Sigma-Aldrich. The lysozyme assay with purified His-tagged proteins was performed photometrically as described previously (Boller et al., 1983; Brunner et al., 1998). Effects of purified His-tagged proteins in 25 mM sodium acetate buffer (pH 5.0) on a growing mycelium of the fungus *Trichoderma viride* (Guangdong Culture Collection Center) were tested at 27°C according to a previously established procedure (Schlumbaum et al., 1986). Under these conditions, the MtNFH1 protein remained stable for at least 20 h as measured with the substrate NodSm-IV(C16:2, S).

Homology Modeling and Substrate-Docking Simulation

The crystal structures of class V chitinases from tobacco (MtChiV; Protein Data Bank [PDB] codes 3ALF and 3ALG) and Arabidopsis (AtChiC; PDB code 3AQU) served as structural templates for homology modeling. The sequence alignment used for modeling was obtained interactively by using the program Swiss-PdbViewer (<http://spdbv.vital-it.ch/>). A model for MtNFH1 was created by using the program MODELER (Fiser and Šali, 2003). The structure of substrate NodSm-V(C16:2, S) was built based on the structure of (GlcNAc)₅ in complex with chitinase B from *Serratia marcescens* (SmChiB; PDB code 1E6N). The substrate NodSm-V(C16:2, S) was then docked manually to the active site of MtChi5, guided by the conserved interactions between the chitinase and the carbohydrate moiety of NodSm-V(C16:2, S). The docked structure of MtChi5 in complex with NodSm-V(C16:2, S) was energy minimized by the program GROMACS (Van Der Spoel et al., 2005) using the topology generated by the program PRODRG (Schüttelkopf and van Aalten, 2004).

Transformation of *M. truncatula* with pCAMBIA-MtNFH1

The transformation of *M. truncatula* ecotype R108-1 was performed by infiltrating leaf discs with *A. tumefaciens* strain EHA105 carrying the RNAi plasmid pCAMBIA-MtNFH1. The transformation and regeneration procedures were based on protocols described by Hoffmann et al. (1997) and Trinh et al. (1998). Briefly, infiltrated leaf explants were placed on petri dishes containing proembryogenic callus induction medium (SHMab) supplemented with acetosyringone. After 3 d of cocultivation with the bacteria, the explants were transferred to new plates containing SHMab medium supplemented with 500 μ g mL⁻¹ cefotaxime and 40 μ g mL⁻¹ hygromycin for selection. Healthy calli emerging from the explants were transferred to plates containing embryogenesis medium (SHM2) supplemented with hygromycin. Plates were changed every 2 weeks until somatic proembryos turned green. The embryos were then kept on SHM2 medium without hygromycin. Plantlets with shoots were transferred onto one-half-strength SHM2 plates and placed vertically to induce root development. Finally, regenerated plantlets were transferred to pots (filled with vermiculite and expanded clay; ratio of 3:1) and used for seed production in a temperature-controlled greenhouse.

GUS Staining

M. truncatula seedlings transformed with the RNAi plasmid pCAMBIA-MtNFH1 were stained for GUS activity with X-Gluc solution [0.5 mg mL⁻¹ 5-bromo-4-chloro-3-indolyl- β -D-glucuronide in 100 mM potassium phosphate buffer, pH 7, 0.1% (v/v) Triton X-100, 10 mM K₃Fe(CN)₆, and 10 mM K₄Fe(CN)₆] at 37°C. Wild-type *M. truncatula* plants served as a control.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers KC833515 (MtNFH1), KC833513 (Mt75352), and KC833514 (MtCRA).

Supplemental Data

- The following materials are available in the online version of this article.
- Supplemental Figure S1.** Unrooted phylogenetic tree of MtNFH1 and 94 predicted plant proteins with related amino acid sequences.
 - Supplemental Figure S2.** Chemical structures of NFs and acylated cleavage products used in this study.
 - Supplemental Figure S3.** Positive-ion MALDI-TOF mass spectra of the cleavage products NodSm-II(C16:2), NodSm-II(C16:2, Ac), and NodSm-III(C16:2) purified by HPLC.
 - Supplemental Figure S4.** The degradation of NodSm-V(C16:2, S) by MtNFH1 is not affected by a 500-fold higher concentration of (GlcNAc)₅.
 - Supplemental Figure S5.** Examples of velocity versus NF substrate concentration curves for MtNFH1.
 - Supplemental Table S1.** MtNFH1, Mt75352, and MtCRA sequences of *M. truncatula* ecotype R108-1 used in this study.

Supplemental Table S2. Positive-ion MALDI-TOF analysis of the cleavage products NodSm-II(C16:2), NodSm-II(C16:2, Ac), and NodSm-III(C16:2).

Supplemental Table S3. K_m and k_{cat} for AtChiC and NtChiV with (GlcNAc)₆ or (GlcNAc)₅ as a substrate.

Supplemental Table S4. Primers used in this study.

Supplemental Table S5. Plasmids constructed in this study.

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