

A novel bacterial β -*N*-acetyl glucosaminidase from *Chitinolyticbacter meiyuanensis* possessing transglycosylation and reverse hydrolysis activities

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Abstract

Background: *N*-acetyl glucosamine (GlcNAc) and *N*-acetyl chitooligosaccharides (*N*-acetyl COSs) exhibit many biological activities, and have been widely used in the pharmaceutical, agriculture, food, and chemical industries. Particularly, higher *N*-acetyl COSs with degree of polymerization from 4 to 7 ((GlcNAc)₄–(GlcNAc)₇) show good antitumor and antimicrobial activity, as well as possessing strong stimulating activity towards natural killer cells. Thus, it is of great significance to discover a β-*N*-acetyl glucosaminidase (NAGase) that can not only produce GlcNAc, but also synthesize *N*-acetyl COSs.

Results: The gene encoding the novel β-*N*-acetyl glucosaminidase, designated *Cm*NAGase, was cloned from *Chitinolyticbacter meiyuanensis* SYBC-H1. The deduced amino acid sequence of *Cm*NAGase contains a glycoside hydrolase family 20 catalytic module that shows low identity (12–35%) with the corresponding domain of most well-characterized NAGases. The *Cm*NAGase gene was highly expressed with an active form in *Escherichia coli* BL21 (DE3) cells. The specific activity of purified *Cm*NAGase toward *p*-nitrophenyl-*N*-acetyl glucosaminide (*p*NP-GlcNAc) was 4,878.6 U/mg of protein. *Cm*NAGase had a molecular mass of 92 kDa, and its optimum activity was at pH 5.4 and 40°C. The V_{max} , K_m , K_{cat} , and K_{cat}/K_m of *Cm*NAGase for *p*NP-GlcNAc were 16,666.67 μmol min⁻¹ mg⁻¹, 0.50 μmol mL⁻¹, 25,555.56 s⁻¹, and 51,111.12 mL μmol⁻¹ s⁻¹, respectively. Analysis of the hydrolysis products of *N*-acetyl COSs and colloidal chitin revealed that *Cm*NAGase is a typical exo-acting NAGase. Particularly, *Cm*NAGase can synthesize higher *N*-acetyl COSs ((GlcNAc)₃–(GlcNAc)₇) from (GlcNAc)₂–(GlcNAc)₆, respectively, showed that it possesses transglycosylation activity. In addition, *Cm*NAGase also has reverse hydrolysis activity toward GlcNAc, synthesizing various linked GlcNAc dimers.

Conclusions: The observations recorded in this study that *Cm*NAGase is a novel NAGase with exo-acting, transglycosylation, and reverse hydrolysis activities, suggests a possible application in the production of GlcNAc or higher *N*-acetyl COSs.

Background

Chitin, a linear polysaccharide of 1, 4-β-linked *N*-acetyl glucosamine (GlcNAc), is the second most abundant renewable source in nature behind cellulose and mainly exists in crustacean shells, fungal cell walls, and insect exoskeletons [1]. Comprehensive utilization of these chitin biomasses may have economic and ecological benefits [2]. However, there is no satisfactory method to utilize them to date. A vast majority of the chitin biomasses are directly disposed of or landfilled without utilization, which leads to serious pollution and wasted resources [3]. Therefore, it would be beneficial to utilize the numerous chitin resources to produce value-added chemicals and materials.

Chitin can be used to produce biomaterials such as films, adhesives, preservative coatings, and antibacterial/anticancer materials [4-6]. Besides the direct utilization of chitin mentioned above, chitin can be used as a substrate for producing nitrogen-containing chemicals like GlcNAc, *N*-acetyl chitooligosaccharides (*N*-acetyl COSs), ethanolamine, N-containing furan derivatives and so on [2].

Among these chemicals, GlcNAc and *N*-acetyl COSs are considered promising platform molecules, and have been widely used in the pharmaceutical, agriculture, food, and chemical industries [7, 8]. Especially, higher *N*-acetyl COSs with the degree of polymerization from 4 to 7 ((GlcNAc)₄–(GlcNAc)₇) exhibit many biological activities. For example, (GlcNAc)₄ was found to have strong stimulating activity towards natural killer cells [9]. (GlcNAc)₅ is an important building block for NOD factor synthesis [10]. (GlcNAc)₆ and (GlcNAc)₇ show antitumor activity against mice sarcoma 180 [11] and antimicrobial activity against fungal pathogens [12].

Chitin is traditionally chemically degraded to GlcNAc and *N*-acetyl COS_s using acid, which leads to toxicity and risks associated with serious pollution during the production process [13]. With increased environmental awareness, increasing attention has been paid to developing enzymatic hydrolysis of chitin using chitinolytic enzymes as catalysts because they are environmentally friendly and result in products with high bioactivity compared to classical chemical routes [14].

Chitinolytic enzymes, the essential enzymes involved in catabolism of chitin, primarily include chitinase (mainly belonging to glycoside hydrolase (GH) families 18 and 19, hydrolyzes chitin to *N*-acetyl COSs) [15], β-*N*-acetyl glucosaminidase (GH families 3, 20, 73, 84, and 85, hydrolyzes *N*-acetyl COSs to GlcNAc) [16, 17], and lytic polysaccharide monooxygenase (Auxiliary Activity (AA) families 10 and 11, cleavage of chitin chains with oxidation to enhance the hydrolysis of chitin) [18]. Of these, β-*N*-acetyl glucosaminidase (NAGase), an indispensable member of the chitinolytic system, has a significant physiological role depending on its origin. With the cooperative action of NAGase and chitinase, chitin can be hydrolyzed into GlcNAc. Moreover, some GH20 NAGases can also be used to synthesize high-value GlcNAc-containing products and *N*-acetyl COSs by transglycosylation and reverse hydrolytic reactions [19-24]. These excellent features make GH20 NAGases receive increased attention. GH20 NAGases can be found in a wide variety of organisms including bacteria, fungal, insects, plants, and animals [24-27]. However, studies about the GH20 NAGases with transglycosylation and reverse hydrolysis activities are mainly derived from fungal sources [28-34]. There are few reports about the bacterial NAGases possessing transglycosylation and reverse hydrolysis activities [21, 35-37].

In our previous study, a chitinolytic bacterium *Chitinolyticbacter meiyuanensis* SYBC-H1 with a good ability to degrade chitin, was isolated from soil [38]. In this study, a gene encoding NAGase was cloned from the SYBC-H1 strain, based on the results of peptide mass fingerprinting and complete genome sequencing, and heterologously expressed in *Escherichia coli* BL21(DE3). The phylogenetic relationships and catalytic characteristics of the purified recombinant NAGase were described. Furthermore, its transglycosylation and reverse hydrolysis activities were also investigated, respectively.

Results And Discussion

Purification of wild-type NAGase from *C. meiyuanensis* SYBC-H1

Purification of the NAGase from *C. meiyuanensis* SYBC-H1 was performed. Following ammonium sulfate precipitation, anion exchange chromatography, and SDS-PAGE, a protein band with NAGase activity was obtained, as shown by zymogram analysis. The band with NAGase activity (named *CmNAGase*) had a molecular mass of between 75 kDa and 100 kDa (Fig. 1).

The protein strip in the stained gel was excised for peptide mass fingerprinting (PMF) analysis using matrix-assisted laser desorption ionization-time-of-flight (MALDI-TOF MS/MS), and the results of PMF were interpreted by referencing the Mascot database [39]. Proteins receiving the highest molecular weight search scores (MOWSE) were selected as the peptide fragments of purified protein. Peptide fragments of purified protein were mainly detected with the amino acid sequences of YDGD TFLARLT LTNH, AMNVR YERLVKAGK, and WNQFANRLGQRELARLDGFLGGYGYRVPV, which showed 100% identity to the peptides from an annotated NAGase in the complete genome of *C. meiyuanensis* SYBC-H1.

Cloning of the *CmNAGase* gene and sequence analysis. The *CmNAGase* gene was cloned. As expected, a PCR product of 2.5 kb was obtained. Analysis of the PCR product showed that the *CmNAGase* gene was 2,508 bp, encoding for a protein of 836 amino acids. The calculated molecular mass of *CmNAGase* was 91.6 kDa, and the isoelectric point (pI) was predicted to be 5.48. The sequence analysis suggested no putative signal peptide in the sequence of *CmNAGase*, which suggested that *CmNAGase* should be a non-secretory protein. The prediction using Gneg-mPLoc 2.0 also showed that the location of *CmNAGase* was in the periplasm. However, *CmNAGase* could be purified from the fermentation broth of the strain SYBC-H1. Furthermore, most reported NAGases are secretory proteins to date [21-23]. These results show that the prediction of *CmNAGase* location may be wrong.

According to the Carbohydrate-Active enZYmes (CAZy) database (<http://www.cazy.org/>), NAGases can be classified as part of the glycoside hydrolase (GH) families 3, 20, 73, 84, and 85 based on amino acid sequence homology. BLASTP analysis showed that *CmNAGase* belonged to GH family 20 (GH20) and shared the highest identity (81.68%) with the GH20 NAGase Ch1 from *Chitiniphilus shinanonensis* (WP_018749679) [40], followed by GH20 NAGase (81.44%) from *Chitiniphilus* sp. HX-2-15 (WP_136772659). However, these coding genes have not been expressed and studied. Among characterized GH20 NAGases, *CmNAGase* showed the highest identity (77.58%) with GH20 NAGase from *Aeromonas* sp. 10S-24 (accession no. BAA92145) [41], following by the GH20 NAGase (34.02%) from *Serratia marcescens* (PDB 1QBA) [42], GH20 NAGase (30.44%) from *Aeromonas caviae* CB101 (accession no. CAH55822) [43], GH20 NAGase Nag2 (30.04%) from *Vibrio harveyi* (PDB 6EZR) [44], GH20 NAGase (29.01%) from *Enterobacter* sp. G-1 (accession no. BAA74506) [45], and GH20 NAGase (24.64%) from *Aeromonas hydrophila* SUWA-9 (accession no. BAF76001) [46], GH20 NAGase Nag1 (12.26%) from *V. harveyi* (accession no. ADJ68332) [44], and GH20 NAGase (12.24%) from *Arthrobacter* sp. TAD20 (accession no. CAB72127) [47]. The putative GH20 NAGases with the highest similarity of *CmNAGase* and verified GH20 NAGases were performed to construct the phylogenetic tree, which also showed that *CmNAGase* exhibited a low sequence identity (12–35%) with most of functionally-characterized bacterial GH20 NAGases (Fig. 2).

GH20 NAGases employ the retaining mechanism for catalysis. The enzymes carry out substrate-assisted catalysis, in which a Glu (E509 in *CmNAGase*) acts as the general acid/base residue for protonation, while Asp (D508 in *CmNAGase*) acts to orient the C2-acetamido group into position for correct nucleophilic attack by a water molecule, and subsequently provides the negatively charged carboxylate groups to stabilize the positively charged oxazolinium ion intermediate [26]. Multiple alignments of the catalytic domain in *CmNAGase* with other GH20 NAGases from different sources indicated the substrate binding residues (R315, H422, V463, Q464, W558, W594, Y621, D623, L624, Y635, W637, W693, and E695) and catalytic residues (D508 and E509) in *CmNAGase*, which are highly conserved among GH20 members (Fig. S1). Moreover, sequence ⁵⁰⁴H/N-X-A/C/G/M-D-E-A/I/L/V⁵¹⁰ in *CmNAGase* is the highly conserved amino acid sequence in the catalytic domain of GH20 NAGases from bacteria, fungi, and archaea [48]. The analysis of secondary structure showed that *CmNAGase* possesses 21 α -helices and 31 β -sheets with the typical TIM-barrel(β/α)₈ fold in the GH20 catalytic domain (Fig. S1), which is consistent with various GH20 NAGases from different sources [49].

Domain structure prediction revealed that *CmNAGase* contains four domains: CHB_HEX domain of residues 7–155 (putative carbohydrate binding domain); Glyco_hydro_20b domain of residues 174–300 (N-terminal domain of the beta-hexosaminidases); Glyco_hydro_20 domain of residues 304–719 (catalytic domain); and CHB_HEX C_1 domain of residues 753–828 (unknown function) (Fig. S2a). As shown in Fig. S2b, the 3D structure of *CmNAGase* was predicted based on the structure model of 1QBA (34.02%) [42]. The active sites (R315, H422, V463, Q464, D508, E509, W558, W594, Y621, D623, L624, Y635, W637, W693, and E695) were also shown in the 3D structure of *CmNAGase* (Fig. S2c), and form an active pocket [48].

Expression of *CmNAGase* gene and purification of recombinant *CmNAGase*

The full-length of *CmNAGase* gene was successfully expressed in *E. coli* BL 21 (DE3) at a high expression level (~50% of total protein). The recombinant *CmNAGase* with a C-terminal His6-tag was purified by Ni-NTA affinity chromatography with a yield of 80.5%. The reason for the high purification yield may be that the C-terminal His6-tag in the recombinant *CmNAGase* was well exposed, which led to the higher affinity with Ni-NTA resin. The specific activity of recombinant *CmNAGase* increased 1.5-fold from 3,156.5 U/mg to 4,878.6 U/mg after purification (Table S2). The SDS-PAGE analysis showed that purified recombinant *CmNAGase* possesses a high purity with an approximate molecular weight of 92 kDa, which agrees with 92,571 kDa calculated from the amino acid sequence containing the His6-tag (Fig. S3).

Effects of pH and temperature on activity and stability of recombinant *CmNAGase*

The pH and temperature profile of *CmNAGase* activity are shown in Fig. 3. Typically, the optimal pH of reported GH20 NAGases are in the range of pH 5.0 to pH 8.0. *CmNAGase* exhibited a high level of activity at pH 4.0–7.0 with the optimal pH of 5.4 (Fig. 3a), which is different from that of NAGases from *Aeromonas* sp. 10S-24 (7.0) [41], *Paenibacillus* sp. TS12 (6.0) [50], *Vibrio harveyi* 650 (7.5) [44], *Enterobacter* sp. G-1 (6.0) [45], *Salmonella enterica* (4.0) [51], *Paraglaciecola hydrolytica* S66 (6.0) [21],

and *Cellulomonas fimi* (7.3–8.7) [52]. In addition, *CmNAGase* presented good activity after being stored at pH 4.0–8.5 for more than 84 h (Fig. 3b), which suggested that the *CmNAGase* possesses a good pH stability compared with other reported NAGases [46, 53, 54].

As shown in Fig. 3c, the effect of temperature on enzymatic activity showed that the optimal temperature of *CmNAGase* was 40°C, which is different from that of NAGases from *Shinella* sp. (50°C) [27], *Trichoderma reesei* (60°C) [53], *P. hydrolytica* S66 (50°C) [21], *Enterobacter* sp. G-1 (45°C) [45], and *S. marcescens* (52°C) [55]. The *CmNAGase* was unstable at temperatures \geq 40°C (Fig. 3d). Half-lives of *CmNAGase* toward 30°C, 35°C, 40°C, and 45°C were 13.0, 9.5, 6.3, and 0.6 h, respectively (Table S3), which were similar with that of GH20 NAGases from *Lactobacillus casei* [56], *Sphingobacterium* sp. [57], and *Aeromonas* sp. 10S-24 [41]. These results suggested that *CmNAGase* is a mesophilic and acidic enzyme.

Effect of metal ions on activity of *CmNAGase*

The effects of metal ions on *CmNAGase* were investigated. All counter-ions of the used metal ions were Cl⁻. As shown in Table 1, EDTA did not inhibit the enzymatic activity, which indicates that *CmNAGase* is not metal-dependent. *CmNAGase* activity is completely inhibited by Zn²⁺, Cu²⁺, and Al³⁺, severely inhibited by Ba²⁺, Fe³⁺, and Cr³⁺. To date, many studies have shown that Zn²⁺, Cu²⁺, Fe³⁺, and Al³⁺ inhibit the activity of NAGases. For example, the GH20 NAGase from *A. caviae* is strongly inhibited by Cu²⁺ and Zn²⁺ [43]; the GH20 NAGase from *Paenibacillus* sp. is strongly inhibited by Zn²⁺ [50], and the GH20 NAGase from *T. reesei* is partially inhibited by Fe³⁺ [53]. Mn²⁺ enhanced the activity of *CmNAGase*, which is different from the GH20 NAGase from *A. caviae* (strongly inhibited by Mn²⁺) [43]. However, the specific activated mechanism of Mn²⁺ is unclear and the in-depth study is needed in the future.

The substrate specificity of *CmNAGase*

The specific activities of the recombinant *CmNAGase* against various substrates were investigated using standard assay conditions. As shown in Table 2, *CmNAGase* can hydrolyze *p*NP-GlcNAc, 4-MU-GlcNAc, and (GlcNAc)₂–(GlcNAc)₆. No activity was observed when *p*NP-glucose, *p*NP-acetyl galactosaminide, and cellobiose were used as the substrates. These results showed that *CmNAGase* represents the typical NAGase activity with strict substrate specificity. The specific activity of *CmNAGase* toward *p*NP-GlcNAc can reach 4,878.6 U/mg, which is higher than most reported NAGases (<2,000 U/mg) [17, 21, 27, 50, 58].

Most of the GH20 NAGases have the highest catalytic efficiency for (GlcNAc)₂ among natural substrates, and do not hydrolyze *N*-acetyl COSs and chitin polymer [49, 58]. However, *CmNAGase* showed good activities toward (GlcNAc)₂–(GlcNAc)₆ with the highest activity for (GlcNAc)₂, followed by (GlcNAc)₃, (GlcNAc)₄, (GlcNAc)₅, and (GlcNAc)₆. These results show that specific activity of *CmNAGase* toward *N*-acetyl COSs decreased when the degree of polymerization increased, which is similar with other reports [59]. The catalytic efficiency of GH20 *LeHex20A* from *Lentinula edodes* for (GlcNAc)₆ was greater than for (GlcNAc)₂ [60]. The GH20 NAGase *VhNag2* from *V. harveyi* showed the highest activity against (GlcNAc)₄,

while the lowest activity was observed with $(\text{GlcNAc})_2$ [61]. These NAGases are different from *CmNAGase*. Furthermore, *CmNAGase* showed some activity (0.02 U/mg) toward colloidal chitin. The result is similar with the GH20 NAGases from *V. harveyi* [61] and the fungal NAGases from *Myceliophthora thermophila* [62], *L. edodes* [60], and *M. anisopliae* [63], which were reported to degrade chitin to some extent without the cooperation of chitinase.

In addition, the kinetic parameters for *CmNAGase* were also measured with *p*NP-GlcNAc as the substrate. The results showed that the V_{max} , K_m , K_{cat} , and K_{cat}/K_m for *CmNAGase* were $16,666.67 \mu\text{mol}\cdot\text{min}^{-1}\cdot\text{mg}^{-1}$, $0.5 \mu\text{mol mL}^{-1}$, $25,555.56 \text{ s}^{-1}$, and $51,111.12 \text{ mL } \mu\text{mol}^{-1} \text{ s}^{-1}$, respectively.

Hydrolysis reaction of recombinant *CmNAGase* toward colloidal chitin.

As shown in Fig. 4a, hydrolysis of colloidal chitin resulted in GlcNAc as the only product, and its concentration increased with the increase of hydrolysis time. Konno et al. [60] reported that the NAGase *LeHex20A* from *L. edodes* hydrolyzes colloidal chitin to various *N*-acetyl COSs at the start of the reaction, and these *N*-acetyl COSs convert to GlcNAc after 3 h, which is different from *CmNAGase*.

Hydrolysis and transglycosylation reactions of recombinant *CmNAGase* toward *N*-acetyl COSs.

To evaluate the hydrolysis and transglycosylation activities of *CmNAGase*, $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_6$ were used as substrates. The overall rates of hydrolysis were in the order: $(\text{GlcNAc})_2 > (\text{GlcNAc})_3 > (\text{GlcNAc})_4 > (\text{GlcNAc})_5 > (\text{GlcNAc})_6$, which is consistent with the results of substrate specificities above. $(\text{GlcNAc})_2$ was degraded by *CmNAGase* to GlcNAc (Fig. 4b). GlcNAc and $(\text{GlcNAc})_2$ are produced from $(\text{GlcNAc})_3$ (Fig. 4c). When using $(\text{GlcNAc})_4$ as a substrate, GlcNAc, $(\text{GlcNAc})_2$, and $(\text{GlcNAc})_3$ were produced (Fig. 4d). GlcNAc, $(\text{GlcNAc})_2$, $(\text{GlcNAc})_3$, and $(\text{GlcNAc})_4$ were obtained from $(\text{GlcNAc})_5$ (Fig. 4e). When using $(\text{GlcNAc})_6$ as the substrate, GlcNAc, $(\text{GlcNAc})_2$, $(\text{GlcNAc})_3$, $(\text{GlcNAc})_4$, and $(\text{GlcNAc})_5$ were found (Fig. 4f). Finally, $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_6$ were both hydrolyzed to GlcNAc as the final product with the increase of hydrolysis time. Similar results were reported for GH20 NAGase (*VhNag2*) from *V. harveyi* [61] and *LeHex20A* from *L. edodes* [60]. However, the hydrolysis rates of *VhNag2* and *LeHex20A* toward *N*-acetyl COSs were in the order: $(\text{GlcNAc})_4 > (\text{GlcNAc})_3 > (\text{GlcNAc})_5 > (\text{GlcNAc})_6 > (\text{GlcNAc})_2$. Based on these results, we conclude that *CmNAGase* is a typical exo-acting NAGase.

In addition, minor $(\text{GlcNAc})_3$, $(\text{GlcNAc})_4$, $(\text{GlcNAc})_5$, $(\text{GlcNAc})_6$, and new peak generated after the peak of $(\text{GlcNAc})_6$ were also produced from $(\text{GlcNAc})_2$, $(\text{GlcNAc})_3$, $(\text{GlcNAc})_4$, $(\text{GlcNAc})_5$, and $(\text{GlcNAc})_6$ in short reaction times, respectively (Fig. 4b–4f). The mass spectrum analysis of the new peak is at *m/z* values of 1440.5756 and 1441.5743, which respectively correspond to $(\text{GlcNAc})_7$ and $(\text{GlcNAc})_7$ with a hydrogen adduct (Fig. S4). These results showed that *CmNAGase* can produce higher *N*-acetyl COSs $(\text{GlcNAc})_3$ – $(\text{GlcNAc})_7$ from $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_6$, respectively. To date, most scholars all believe that production of $(\text{GlcNAc})_{n+1}$ from $(\text{GlcNAc})_n$ ($n=2-7$) is a transglycosylation reaction [28, 29, 34]. However, it may also be a reverse hydrolysis reaction. For example, $(\text{GlcNAc})_2$ and $(\text{GlcNAc})_n$ may form $(\text{GlcNAc})_{n+1}$ by

transglycosylation. Meanwhile, GlcNAc and $(\text{GlcNAc})_n$ could also form $(\text{GlcNAc})_{n+1}$ through reverse hydrolysis. It is very hard to tell the two reactions apart in this setup. Thus, we call this reaction transglycosylation for the time being.

To date, some chitinases have been used to synthesize higher *N*-acetyl COSs from shorter *N*-acetyl COSs substrates via transglycosylation activity. For example, a chitinase from *C. shinanonensis* generated $(\text{GlcNAc})_5$ and $(\text{GlcNAc})_6$ when incubated with $(\text{GlcNAc})_4$ [64]. A chitinase from *T. reesei* KDR-11 was shown to convert $(\text{GlcNAc})_4$ into $(\text{GlcNAc})_6$ [65]. An endochitinase of *Flavobacterium johnsoniae* synthesized $(\text{GlcNAc})_6$ – $(\text{GlcNAc})_8$ and $(\text{GlcNAc})_7$ – $(\text{GlcNAc})_9$ from $(\text{GlcNAc})_5$ and $(\text{GlcNAc})_6$, respectively [66]. A chitinase from *Microbulbifer thermotolerans* DAU221 produced $(\text{GlcNAc})_4$ from $(\text{GlcNAc})_3$ [67].

For GH20 NAGases, production of higher *N*-acetyl COSs from shorter *N*-acetyl COSs were often catalyzed by an auto-condensation reaction (a special case of transglycosylation, which involves only one substrate (acts both donor and acceptor) [24]. For example, a GH20 NAGase from *A. oryzae* was used to catalyze the formation of $(\text{GlcNAc})_3$ and $(\text{GlcNAc})_4$ from $(\text{GlcNAc})_2$ [29]. Singh et al. [28] reported that the GH20 NAGase from *A. oryzae* produce $(\text{GlcNAc})_4$ – $(\text{GlcNAc})_6$ and $(\text{GlcNAc})_5$ – $(\text{GlcNAc})_6$ from $(\text{GlcNAc})_3$ and $(\text{GlcNAc})_4$, respectively. GH20 NAGases *SmHex* from *S.marcescens* YS-1 [68] and *NoHex* from *N.orientalis* IFO12806 [69] were shown to convert $(\text{GlcNAc})_2$ to $(\text{GlcNAc})_3$. GH20 NAGases *AoHex* from *A. oryzae* CCF1066 produce the mixture of $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_8$ from the mixture of $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_7$ [34]. In comparison with these reports, *CmNAGase* is a novel bacteria-derived NAGase, which possesses transglycosylation activity toward $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_6$.

In addition, the products $(\text{GlcNAc})_3$ – $(\text{GlcNAc})_7$ from transglycosylation disappeared soon with the increase of reaction time (Fig. 4b-4f). This phenomenon may be because the NAGase activity of *CmNAGase* outweighs its transglycosylation activity, which leads to the transient existence of $(\text{GlcNAc})_3$ – $(\text{GlcNAc})_7$.

Reverse hydrolysis activity of *CmNAGase* toward GlcNAc

In view of the transglycosylation activity toward $(\text{GlcNAc})_2$ – $(\text{GlcNAc})_6$, the reverse hydrolysis activity of *CmNAGase* was also investigated using GlcNAc as the substrate. As shown in Fig. 5, two new peaks at 8.2 min (peak 1) and 10.9 min (peak 2) were detected by HPLC. Of these, peak 1 was $(\text{GlcNAc})_2$ compared with the standard of $(\text{GlcNAc})_2$. However, the retention time of peak 2 was between that of $(\text{GlcNAc})_2$ (8.2 min) and $(\text{GlcNAc})_3$ (12.3 min). To further identify peak 2, mass spectrum analysis was conducted. The *m/z* value of peak 2 was at 447.1586, which corresponds to $(\text{GlcNAc})_2$ (425.1766 Da) with a sodium adduct (22.9898 Da) (Fig. S5). The result showed that peak 2 was also GlcNAc dimer.

The reverse hydrolysis reaction of GH20 NAGases toward GlcNAc can form many connection configurations of the GlcNAc dimer, for example by $\beta(1\rightarrow3)$, $\beta(1\rightarrow4)$, and $\beta(1\rightarrow6)$ glycosidic bonds, which is mainly determined by the regioselectivity of glycosidase [30]. To date, studies on the synthesis of

GlcNAc dimer from GlcNAc were mainly reported by Rauvolfová et al, which produce β -GlcNAc-(1 \rightarrow 3)-GlcNAc, β -GlcNAc-(1 \rightarrow 4)-GlcNAc, and β -GlcNAc-(1 \rightarrow 6)-GlcNAc using the library of fungal GH20 NAGases [30]. For example, the GH20 NAGases from *Acremonium persicinum* CCF 1850, *A. oryzae* CCF 1066, *Aspergillus flavipes* CCF 2026, *A. flavus* CCF 1129, *P. oxalicum* CCF 2315, and *A. terreus* CCF 2539, which only produced β -GlcNAc-(1 \rightarrow 6)-GlcNAc from GlcNAc [30, 31]. The GH20 NAGases from *P. funiculosum* CCF 1994, *P. funiculosum* CCF 2325, *P. chrysogenum* CCF 1269, and *Trichoderma harzianum* CCF 2687 synthesized β -GlcNAc-(1 \rightarrow 3)-GlcNAc, β -GlcNAc-(1 \rightarrow 4)-GlcNAc, and β -GlcNAc-(1 \rightarrow 6)-GlcNAc from GlcNAc [30]. The GH20 NAGases from *A. fumigatus* CCF 1059, *P. pittii* CCF 2277, and *A. sojae* CCF 3060 synthesized β -GlcNAc-(1 \rightarrow 4)-GlcNAc and β -GlcNAc-(1 \rightarrow 6)-GlcNAc from GlcNAc. Among these fungal NAGases, β -GlcNAc-(1 \rightarrow 6)-GlcNAc is always the main product. Thus, the peak 1 in Fig. 5 was β -GlcNAc-(1 \rightarrow 4)-GlcNAc ((GlcNAc)₂), and main peak 2 should be β -GlcNAc-(1 \rightarrow 6)-GlcNAc. These results suggested that *Cm*NAGase has reverse hydrolysis activity toward GlcNAc.

Conclusions

This study reports the isolation, cloning, and recombinant expression of the gene encoding *Cm*NAGase from *C. meiyuanensis* SYBC-H1. *Cm*NAGase contains a GH20 family catalytic module and exhibits low similarity with reported GH20 NAGases. Analysis of the hydrolysis products from *N*-acetyl COSs and colloidal chitin revealed that *Cm*NAGase exhibited exo-acting activity. Interestingly, *Cm*NAGase possesses transglycosylation activity toward (GlcNAc)₂-(GlcNAc)₆, which respectively leads to synthesis of (GlcNAc)₃-(GlcNAc)₇. In addition, *Cm*NAGase also have reverse hydrolysis activity toward GlcNAc, which can produce its dimers with different linked. This is first report of a bacterial NAGase, which can produce GlcNAc dimers from GlcNAc via reverse hydrolysis activity.

Methods

Chemicals

Chitin, 4-methylumbelliferyl *N*-acetyl glucosaminide (4-MU-GlcNAc), *p*-nitrophenyl *N*-acetyl glucosaminide (*p*NP-GlcNAc), *p*NP-glucose, and *p*NP-acetyl galactosaminide were purchased from Aladdin reagent Co., Ltd (Shanghai, China). The standards of *N*-acetyl chitooligosaccharides (*N*-acetyl COSs) (purity: \geq 95%) with degree of polymerization between 2–6 were acquired from Qingdao Bozhi Biotechnology Co., Ltd (Qingdao, China). Peptone and yeast extract were purchased from the Oxoid Co., Ltd. (Beijing, China). All molecular reagents were purchased from TaKaRa (Dalian, China). Colloidal chitin was prepared as described by Gao et al. [13]. Other chemicals and solvents used in this study were purchased from local suppliers and were of analytical grade.

Strains, culture conditions, and plasmids

The *C. meiyuanensis* strain SYBC-H1 (ATCC BAA-2140) used in this study was isolated previously [38]. SYBC-H1 was cultivated according to our previous study [70]. The supernatant was collected as crude

enzyme by centrifugation at $6,000 \times g$ at 4°C and used for NAGase purification.

The strains, plasmids, and primers used in this study are listed in Table S1. *E. coli* strains were routinely cultivated aerobically at 37°C in LB medium (10 g/L tryptone, 5 g/L yeast extract, and 5 g/L NaCl). *E. coli* transformants were grown in LB medium or on agar plates containing 50 $\mu\text{g}/\text{mL}$ kanamycin.

Purification of wild-type NAGase from *C. meiyuanensis* SYBC-H1

NAGase was purified by saturation with ammonium sulfate, followed by anion exchange chromatography, and all purification procedures were carried out at 4°C . The supernatant of the culture was used as a crude enzyme preparation and then fractionated at 40% to 60% saturation with ammonium sulfate. The precipitate was centrifuged at 12,000 g for 30 min and dissolved in a suitable volume of 50 mM PBS (pH 7.0). The enzyme obtained in the previous step was further purified using a fast protein liquid chromatography (FPLC) system (AKTA Pure 150; GE healthcare Co., Fairfield, USA) with a DEAE SepharoseTM anion exchange column. The column was equilibrated with 50 mM Tris-HCl at pH 8.0, then protein was separated by gradient elution with NaCl solutions from 0.05 M to 0.5 M. The purified NAGase was concentrated and collected using an ultrafiltration tube (10 kDa, Millipore, USA) at 4°C . Then the purified enzyme was analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) with a 3% stacking gel and a 10% separating gel, according to the method described by Laemmli [71].

After electrophoresis, the gel was sliced vertically into two parts for staining and zymogram analysis. One part was stained using 0.1% Coomassie brilliant blue R-250 and then decolorized with a mixture of 10% acetic acid, 30% methanol, and 60% water. The other part was incubated with 2.5% (vol/vol) Triton X-100 for 30 min twice to refold, then was sprayed with 50 mM PBS (pH 7.0) containing 1 mM 4-MU-GlcNAc and incubated at 37°C for 30 min; the NAGase strip became visible as fluorescence at 340 nm. Two parts were compared to determine the position of the *Cm*NAGase, and the corresponding band was sliced for peptide mass fingerprinting (PMF) analysis.

Peptide mass fingerprinting of the enzyme

The gel sliced above was analyzed using the electrospray ionization quadrupole time-of-flight mass spectrometer (ESI-Q-TOF MS/MS) technique (PROTECH, Inc., Suzhou, China). These masses were then compared to theoretical mass values in the Mascot website databases (<http://www.matrixscience.com>) to reveal the amino acid sequences of the peptide fragments. The peptide sequence was then aligned with the genome of *C. meiyuanensis* SYBC-H1 (GenBank accession number, CP041335) to find the NAGase and its coding gene.

Cloning of the *Cm*NAGase gene and sequence analysis

The genomic DNA of *C. meiyuanensis* SYBC-H1 was used as the template for polymerase chain reaction (PCR) amplification. According to the results of PMF and the complete genome of *C. meiyuanensis* SYBC-

H1, the coding region of the *CmNAGase* gene was amplified by PCR with the primer pair *CmNAGase*F-5'-GAATTCATATGATGAGCCGTCGCCGGATC-3' and *CmNAGase*R-5'-TCCGCTCGAGTCAGGCGCCACCTGCACCG-3'. The PCR conditions were as follows: 5 min at 94°C, followed by 30 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 10 min. The amplified PCR product was purified by gel electrophoresis, digested with restriction enzymes *Nde*I and *Xho*I, and then ligated into pMD19-T Simple vector and sequenced by Invitrogen Corporation (Shanghai, China). The positive recombinant plasmids were digested with *Nde*I and *Xho*I, and the gene was inserted into the pET-28a(+) vector expression plasmid with a C-terminal His6-tag to generate the pET-28a(+)-*CmNAGase*.

Nucleotide and amino acid sequences were analyzed using Snap Gene™ 1.1.3 software and the ExpASy ProtParam tool (<http://web.expasy.org/protparam/>) [72]. The conserved domains and the GH family classification were identified via the SMART website (<http://smart.embl-heidelberg.de/>) [73]. The DNA and protein sequence alignments were performed via the NCBI server with the programs BLASTN and BLASTP (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>) [74], respectively. Phylogenetic trees were inferred using neighbor-joining algorithm in MEGA 7.0 software and assessed using 1,000 bootstrap replications. The presence of a signal peptide and enzyme location were analyzed using the SignalP 5.0 server (<http://www.cbs.dtu.dk/services/SignalP/>) [75] and Gneg-mPLoc server v.2.0 (<http://www.csbio.sjtu.edu.cn/bioinf/Gneg-multi/>) [76]. Protein homologous sequences alignment was carried out using ClustalX 2.1 software and ESPript 3.0 (<http://esprict.ibcp.fr/ESPript/cgi-bin/ESPript.cgi>) [77]. Three-dimensional (3D) structure of *CmNAGase* was predicted with RaptorX (<http://raptorx.uchicago.edu/StructPredV2/predict/>) [78].

Expression of the *CmNAGase* gene in *E. coli* BL21(DE3) and purification of the recombinant enzyme

The recombinant plasmid pET-28a(+)-*CmNAGase* above was transformed into competent *E. coli* BL21(DE3) for protein expression. The *E. coli* BL21(DE3) harboring the pET-28a(+)-*CmNAGase* plasmid were cultured in LB medium (containing 50 µg/mL kanamycin) at 37°C in a shaker with a rotation speed of 200 rpm. When the optical density (OD₆₀₀) of the culture medium reached 0.6–0.8, isopropyl β-D-thiogalactoside (IPTG) was added at a final concentration of 1 mM for protein induction, and the culture was further grown at 25°C for 12 h.

The cells were harvested by centrifugation at 6000 *g* and 4°C for 10 min, after which the cells were re-suspended with His6-tag binding buffer (20 mM Tris-HCl, 500 mM NaCl, 50 mM imidazole [pH 7.0]) and lysed by JY92-IIN ultrasonication (Ningbo Xinzhi Biotechnology, Ltd., Ningbo, China). Cell debris was removed by centrifugation at 6000 *g* for 10 min at 4°C and the supernatant was retained as crude enzyme. The recombinant *CmNAGase* were purified using a FPLC system (AKTA Pure 150; GE healthcare Co., Fairfield, USA) with a Ni-nitrilotriacetic acid affinity chromatography (Ni-NTA) column (His Trap™ FF 5 mL). The target protein was eluted with elution buffer (20 mM Tris-HCl, 500 mM NaCl, 250 mM imidazole [pH 7.0]). The eluted fractions were passed through an ultrafiltration tube of 10 kDa (Millipore, USA) to remove the imidazole with sodium phosphate buffer (pH7.0) and concentrate the enzyme solution.

Determination of protein concentration and molecular weight

Concentration of protein was quantified using the Bradford method [79]. Bovine serum albumin (BSA) was used to construct a standard calibration curve.

Reductive SDS-PAGE with a 3% stacking gel and 10% separating gel was performed to determine the molecular weight of purified recombinant protein according to purification part of wild-type NAGase above. A premixed protein marker (Takara Biotechnology Co., Ltd., Nanjing, China) containing 180-, 140-, 100-, 75-, 60-, 45-, 35-, 25-, 15-, and 10-kDa bands was used as the molecular mass standard.

Determination of enzymatic activity

The NAGase activity for *CmNAGase* used *p*NP-GlcNAc as the substrate [80]. A total of 20 μ L of the enzyme solution (0.1 g/L) was added to 0.98 mL *p*NP-GlcNAc (0.25 mM) in 50 mM sodium citrate buffer (pH 5.4) and incubated at 40°C for 10 min. The reaction was terminated by adding 2 mL NaOH (0.5 M). The absorbance was measured at 405 nm to determine the amount of *p*NP produced using a standard curve. One unit of NAGase activity was defined as the amount of enzyme required to release 1 μ mol *p*NP from the substrate per minute at 40°C.

Characterization of recombinant *CmNAGase*

With 1 mM *p*NP-GlcNAc as the substrate, the optimum pH for activity of *CmNAGase* was determined using different buffers: 50 mM citrate buffer (pH 3.0–6.0), 50 mM phosphate buffer (pH 6.0–7.5), and Tris-HCl buffer (pH 7.0–9.0) at 40°C for 30 min. To measure the pH stability, enzyme was incubated at 35°C for 96 h in the different buffers and the residual activities were determined against 1 mM *p*NP-GlcNAc.

The optimum temperature of *CmNAGase* activity was measured, and the reaction solutions were incubated at temperatures that ranged from 25°C to 55°C for 30 min. Enzyme thermostability was determined by measuring the residual activities after pre-incubation of the purified enzyme in 50 mM sodium citrate buffer (pH 5.4) at 20°C–40°C without substrate for 12 h. The residual activities were performed at pH 5.4 and 40°C according to enzymatic activity assay above.

The effects of metal ions on the activity were also determined. Purified recombinant *CmNAGase* was treated with 10 mM EDTA for 4 h at 4°C and then dialyzed against 50 mM Tris-HCl buffer (pH 7.0) to remove the EDTA. For reactivation, the metal-free enzyme was incubated with various metal salts containing Cr^{3+} (CrCl_3), Fe^{3+} (FeCl_3), Fe^{2+} (FeCl_2), Ca^{2+} (CaCl_2), Cu^{2+} (CuCl_2), Mg^{2+} (MgCl_2), Zn^{2+} (ZnCl_2), Mn^{2+} (MnCl_2), Ni^{2+} (NiCl_2), Co^{2+} (CoCl_2), K^+ (KCl), or Na^+ (NaCl) at final concentrations of 10 mM for 30 min, and the residual activities were then measured with *p*NP-GlcNAc at 40°C and pH 7.0 (Tris-HCl buffer) for 10 min. The activity without addition of metal ions was used as the control (100%).

The substrate specificity of *CmNAGase* was determined by measuring the activity of enzyme toward colloidal chitin, *N*-acetyl COSs ((GlcNAc)₂-(GlcNAc)₆), cellobiose, 4-MU-GlcNAc, and *p*NP-glycosides as substrates) at a concentration of 10 mg/mL. Reaction mixture (1 mL) containing enzyme (3 µg) and various substrates (10 g/L) was incubated at 40°C and pH 5.4 for 10 min. The amount of reducing sugars from colloidal chitin and (GlcNAc)₂-(GlcNAc)₆ were quantified with HPLC. The amount of *p*NP released in the reaction mixture was determined by measuring the absorbance at 405 nm. Hydrolytic activity of the enzyme against *N*-acetyl COSs was assayed by measuring the amount of GlcNAc released during the enzymatic reaction. After incubation, the enzymatic reaction was stopped by heating the mixture in a boiling water bath for 5 min. One unit of enzyme activity was defined as the amount of enzyme required to liberate 1 µmol of *p*NP or GlcNAc per minute under the assay conditions.

Kinetics experiments were performed using *p*NP-GlcNAc as the substrate. The initial velocities were determined by incubating 17 ng purified *CmNAGase* with *p*NP-GlcNAc concentrations ranging from 50 µM to 1000 µM at 40°C in a 1 mL reaction system (50 mM sodium citrate buffer, pH 5.4) for 5 min. The values of V_{max} , K_m , and K_{cat} were estimated by linear regression from double-reciprocal plots according to the method of Lineweaver [81].

Hydrolysis reaction of the recombinant *CmNAGase* toward colloidal chitin

A 1 mL reaction system (50 mM sodium citrate buffer, pH 5.4) containing colloidal chitin (10 g/L) and purified *CmNAGase* (3 µg) was conducted at 35°C for various time intervals. Boiling (5 min) was used to stop the reaction.

Hydrolysis and transglycosylation reactions of the recombinant *CmNAGase* toward *N*-acetyl COSs

A 100 µL volume (50 mM sodium citrate buffer, pH 5.4) with 10 g/L *N*-acetyl COSs ((GlcNAc)₂-(GlcNAc)₆) and 0.3 µg purified *CmNAGase* in a 100 µL volume at 35°C for various time intervals. The enzyme reactions were stopped by boiling at 100°C for 5 min.

Reverse hydrolysis reaction of the recombinant *CmNAGase* toward GlcNAc

The determination of reverse hydrolysis activity was performed with 10 g/L GlcNAc and 0.3 µg purified *CmNAGase* in a 100 µL volume at pH 5.4 and 35°C for 1 h. The reactions were stopped by boiling at 100°C for 5 min.

Analysis of products

The resulting reaction products were analyzed with an Agilent 1260 series HPLC system according to our previous report [80]. The molecular mass of the product was analyzed using electrospray ionization mass spectrometry (ESI-MS, API 2000) with the ESI positive mode. The quadrupole scan mode was used under a capillary voltage of 2.8 kV, cone voltage of 30 V, desolvation gas temperature of 350°C and source temperature of 120°C.

Nucleotide sequence accession number

The sequence for the gene encoding *CmNAGase* cloned from strain SYBC-H1 was deposited in GenBank under accession number no. MN548778.

Abbreviations

GlcNAc, *N*-acetyl glucosamine; NAGase, β -*N*-acetyl glucosaminidase; *N*-acetyl COSs, *N*-acetyl chitooligosaccharides; GH, glycoside hydrolase; 4-MU-GlcNAc, 4-methylumbelliferyl *N*-acetyl glucosaminide, *p*NP-GlcNAc, *p*-nitrophenyl *N*-acetyl glucosaminide; PMF, peptide mass fingerprinting; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; PCR, polymerase chain reaction; FPLC, fast protein liquid chromatography; MOWSE, molecular weight search scores; CAZy, Carbohydrate-Active enZymes.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

All authors have seen and approved the manuscript before submission to *Biotechnology for Biofuels*.

Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare no conflicts of interest.

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Authors' contributions

AZ, KC, and PO designed the work. XM, GW, NZ, YW, and JC constructed the clone of *CmNAGase* gene, purified the recombinant protein, performed enzyme characterization and HPLC studies. AZ conducted

bioinformatic studies. The data was analyzed by AZ, XM, NZ, and JC. The manuscript was written by AZ by taking critical inputs from KC.

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Tables

Table 1 Effect of different metal ions on *CmNAGase*

| Metal ions | Chemicals | Relative activity (%) |
|------------------|-------------------|-----------------------|
| No addition | - | 100 |
| EDTA | EDTA | 101.4 ± 3.2 |
| Na ⁺ | NaCl | 96.8 ± 6.7 |
| K ⁺ | KCl | 101.9 ± 7.8 |
| Fe ²⁺ | FeCl ₂ | 97.6 ± 4.2 |
| Ca ²⁺ | CaCl ₂ | 93.9 ± 6.6 |
| Cu ²⁺ | CuCl ₂ | 0 |
| Ni ²⁺ | NiCl ₂ | 106.3 ± 4.9 |
| Zn ²⁺ | ZnCl ₂ | 0 |
| Mg ²⁺ | MgCl ₂ | 83.3 ± 6.3 |
| Mn ²⁺ | MnCl ₂ | 116.3 ± 3.5 |
| Co ²⁺ | CoCl ₂ | 103.1 ± 3.8 |
| Ba ²⁺ | BaCl ₂ | 34.7 ± 4.6 |
| Al ³⁺ | AlCl ₃ | 0 |
| Fe ³⁺ | FeCl ₃ | 21.3 ± 1.1 |
| Cr ³⁺ | CrCl ₃ | 4.3 ± 2.5 |

Samples were preincubated with various metal ions (10 mM) at pH 7.0 (Tris-HCl buffer) and 4°C for 30 min. The remaining activity was measured with *p*NP-GlcNAc at pH 7.0 (Tris-HCl buffer) and 40°C for 10 min. Activity in the absence of any additives was taken as 100%.

Table 2 Substrate specificity of *Cm*NAGase

| Substrates ^a | Specific activity (U/mg of protein) |
|-----------------------------------|-------------------------------------|
| Colloidal chitin | 0.02 ± 0.001 |
| <i>p</i> NP-GlcNAc | 4,878.6 ± 200.8 |
| <i>p</i> NP-Glucoside | - |
| <i>p</i> NP-Acetylgalactosaminide | - |
| 4-MU-GlcNAc | 4,206.1 ± 283.8 |
| Cellobiose | - |
| (GlcNAc) ₂ | 5,305.4 ± 125.6 |
| (GlcNAc) ₃ | 3,132.8 ± 280.5 |
| (GlcNAc) ₄ | 1,409.4 ± 67.3 |
| (GlcNAc) ₅ | 1,116.2 ± 45.1 |
| (GlcNAc) ₆ | 473.7 ± 36.8 |

-: Activity was not detected.

^a Reaction mixture (1 mL) containing enzyme (3 µg) and substrates (10 g/L) was incubated at pH 5.4 and 40°C for 10 min.

Figures

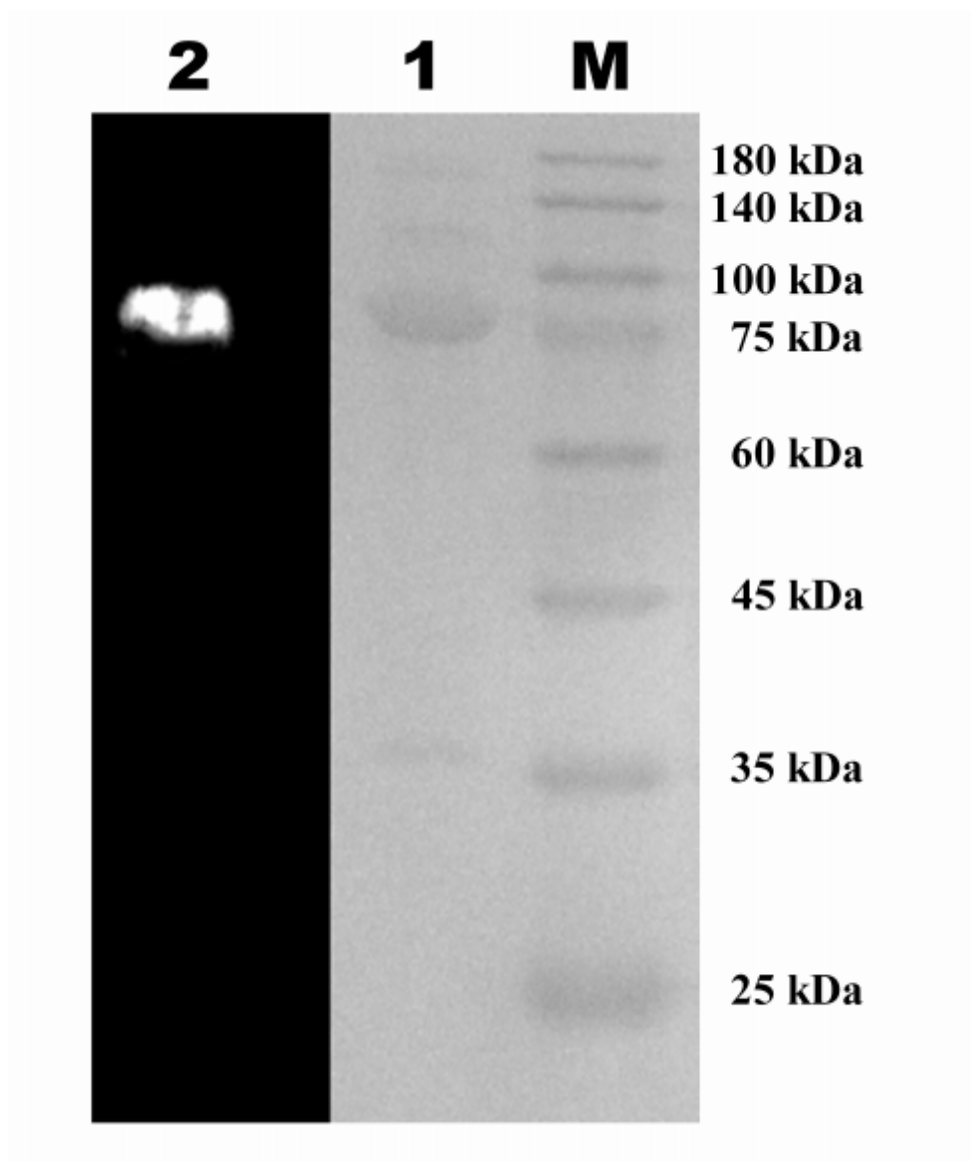


Figure 1

Zymogram analysis of the purified CmNAGase. The purified enzyme was analyzed by SDS-PAGE using a 3% stacking gel and a 10% separating gel with addition of SDS. Lane M, protein molecular mass makers; lane 1, purified CmNAGase (20 µg) visualized by Coomassie brilliant blue staining; lane 2, Purified CmNAGase staining with 4-MU-GlcNAc and visualized at 340 nm.

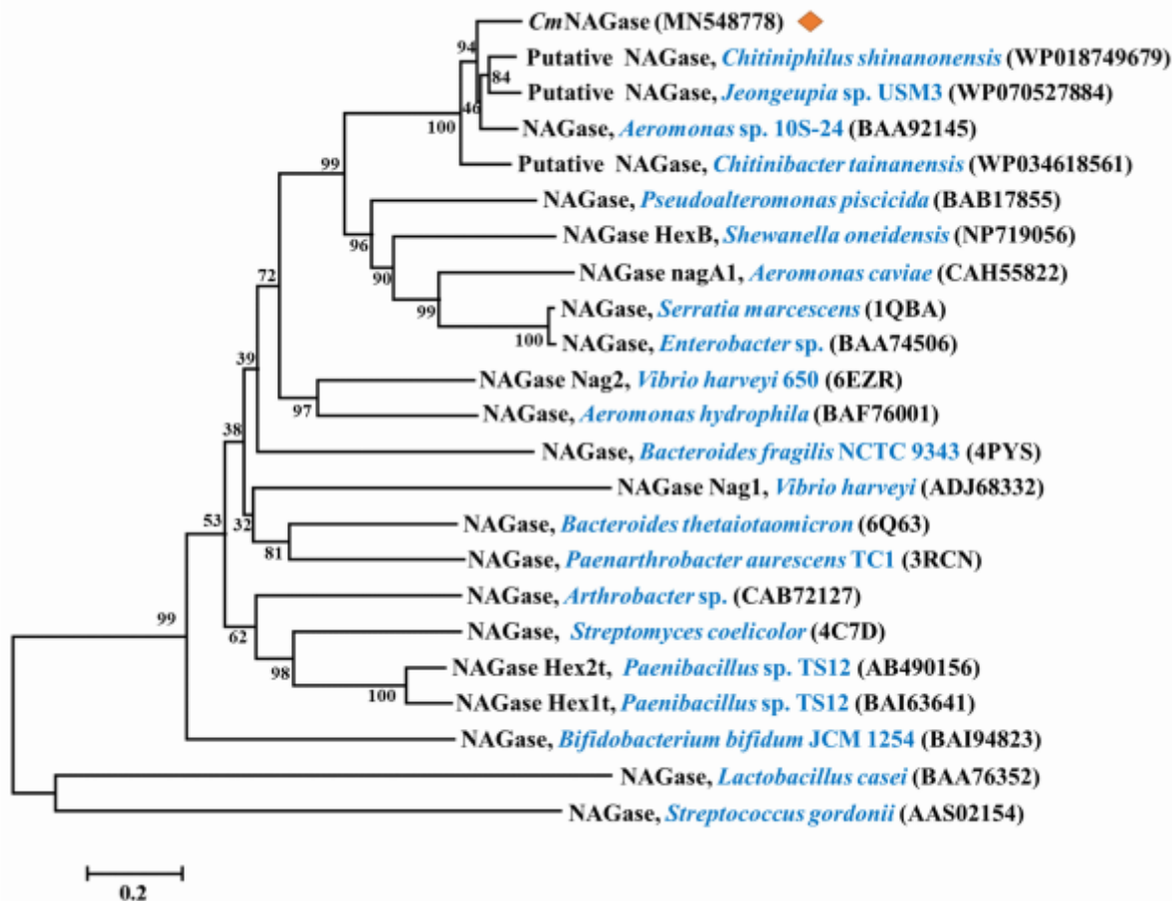


Figure 2

Phylogenetic relationships between CmNAGase and other bacterial NAGases from the glycoside hydrolase (GH) family 20. The phylogenetic tree was constructed by the neighbor-joining algorithm based on the amino acid sequence alignment in MEGA 7.0. The amino acid sequence of CmNAGase was aligned with those of the following proteins: GH20 NAGases from *Chitiniphilus shinanonensis* (WP_018749679), *Jeongeupia* sp. USM3 (WP_070527884), *Aeromonas* sp. 10S-24 (BAA92145), *Chitinibacter tainanensis* (WP_034618561), *Pseudoalteromonas piscicida* (BAB17855), *Shewanella oneidensis* (HexB, NP719056), *Aeromonas caviae* (nagA1, CAH55822), *Serratia marcescens* (PDB 1QBA), *Enterobacter* sp. (BAA74506), *Vibrio harveyi* 650 (Nag2, PDB 6EZR), *Aeromonas hydrophila* (BAF76001), *Bacteroides fragilis* NCTC 9343 (PDB 4PYS), *V. harveyi* (Nag1, ADJ68332), *Bacteroides thetaiotaomicron* (PDB 6Q63), *Paenarthrobacter aurescens* TC1 (PDB 3RCN), *Arthrobacter* sp. (CAB72127), *Streptomyces coelicolor* (4C7D), *Paenibacillus* sp. TS12 (Hex2t, AB490156), *Paenibacillus* sp. TS12 (Hex1t, BAI63641), *Bifidobacterium bifidum* JCM 1254 (BAI94823), *Lactobacillus casei* (BAA76352), and *Streptococcus gordonii* (AAS02154).

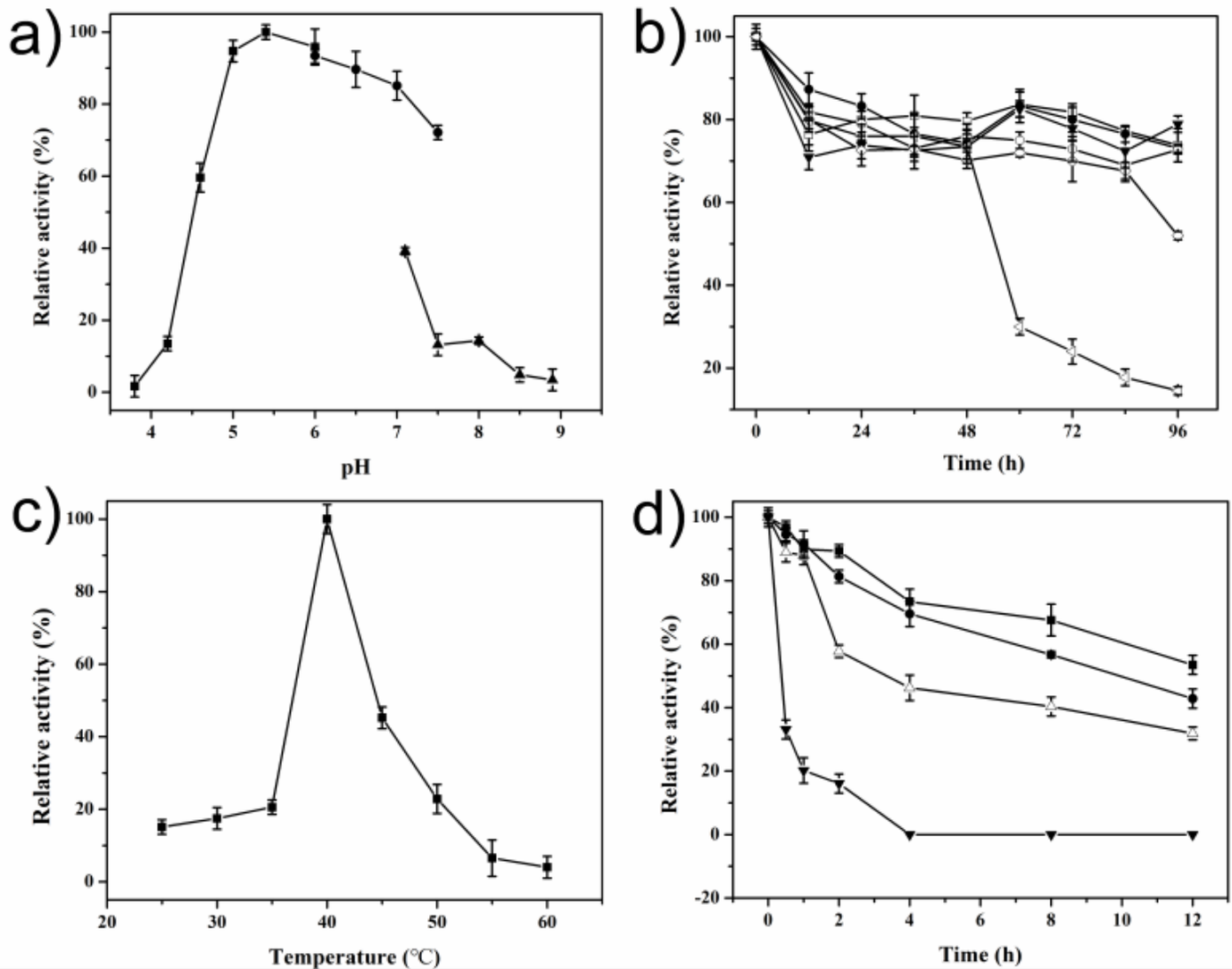


Figure 3

Effect of temperature and pH on CmNAGase. a) The optimal pH of CmNAGase. The optimal pH was determined in 50 mM solutions of various buffers within the pH range of 3.5–10.0. (■, citrate buffer (pH 3.0–6.0); ●, phosphate buffer (pH 6.0–7.5); ▼, Tris-HCl buffer (pH 7.0–9.0)). b) The stability of CmNAGase at various pH values. To determine pH stability, the enzyme was incubated at 35°C for 96 h with various pH buffers, and the residual activities were measured. (▼, 4.7 (citrate buffer); □, 5.4 (citrate buffer); ☒, 6.0 (citrate buffer); ●, 7.0 (phosphate buffer); ●, 8.0 (Tris-HCl buffer); ◁, 9.0 (Tris-HCl buffer)). c) The optimal temperature of CmNAGase. The temperature optimum was determined at different temperatures (25–60°C) in 50 mM sodium citrate (pH 5.4) of the recombinant CmNAGase. d) The stability of CmNAGase at different temperatures. To determine the thermostability, the enzyme was treated in 50 mM sodium citrate (pH 5.4) for 12 h at different temperatures, the residual activity was measured at pH 5.4 and 40°C. (■, 30°C; ●, 35°C; △, 40°C; ▼, 45°C).

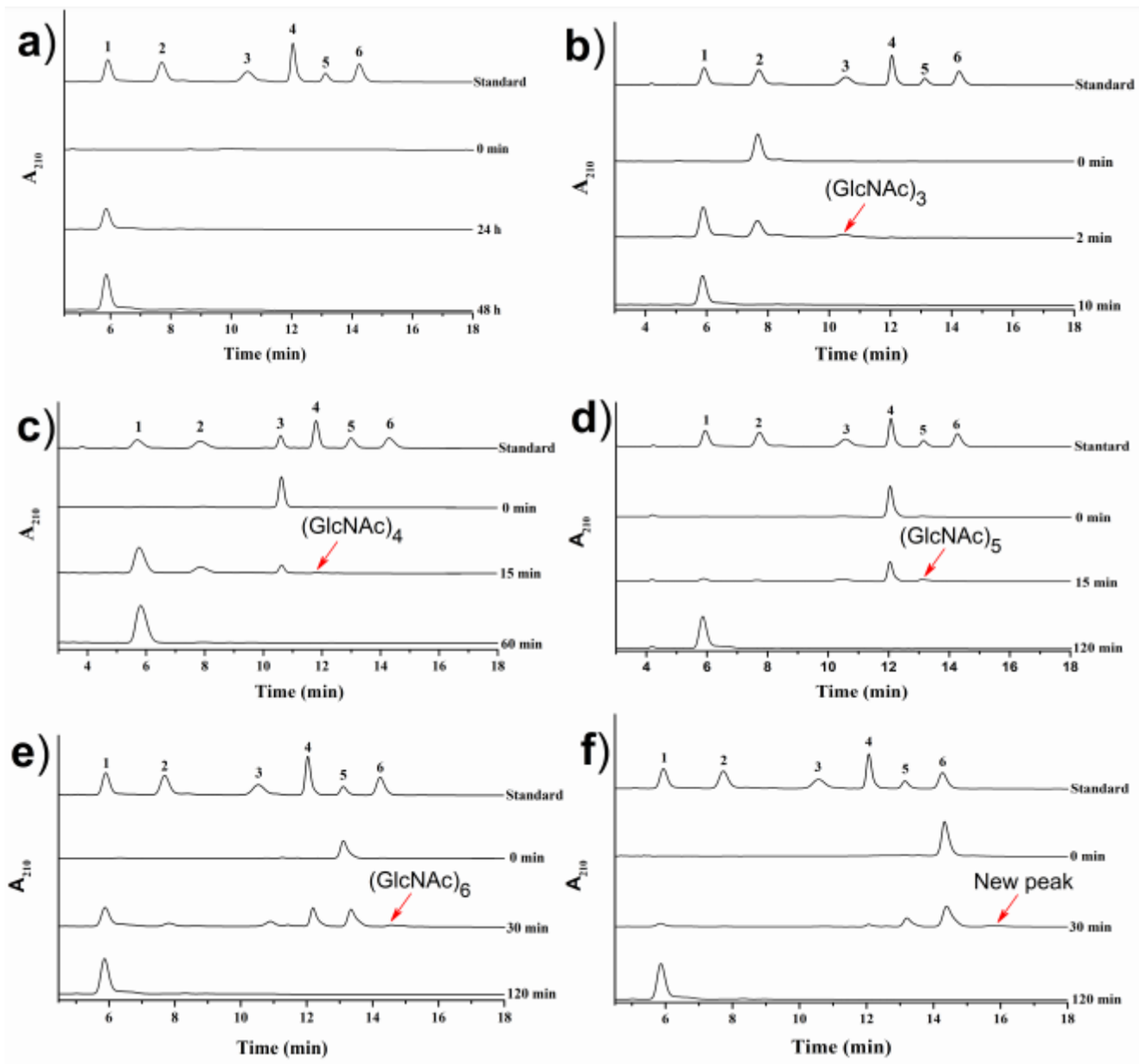


Figure 4

HPLC analysis of the hydrolysate of colloidal chitin and N-acetyl COSs by CmNAGase. a) Colloidal chitin and (b–e) (GlcNAc)₂–(GlcNAc)₆ were incubated with CmNAGase (0.3 μg) in 100 μL volume at pH 5.4 and 35°C for different time intervals. 1–6: Standards of GlcNAc, (GlcNAc)₂, (GlcNAc)₃, (GlcNAc)₄, (GlcNAc)₅, and (GlcNAc)₆.

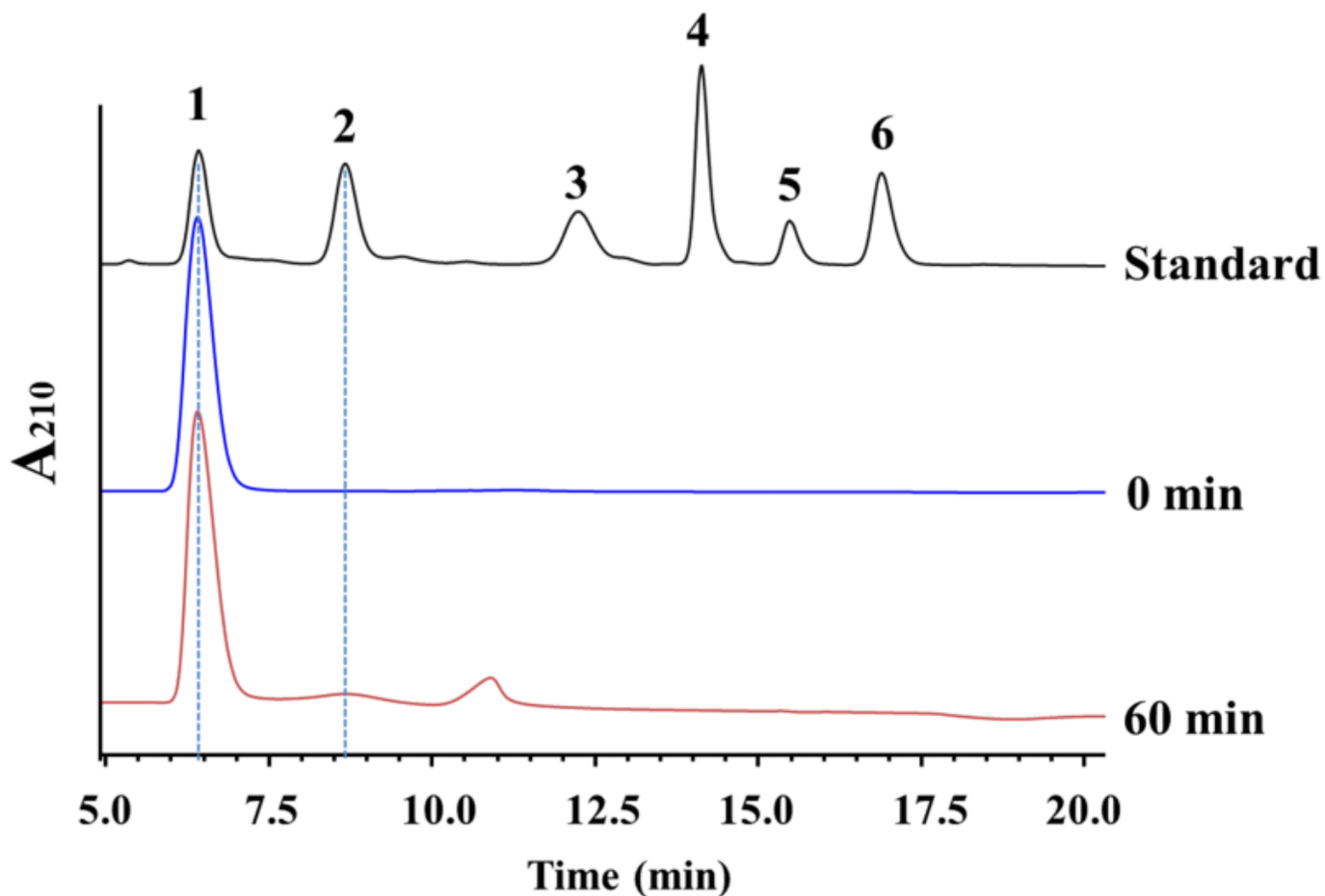


Figure 5

HPLC analysis of the products from GlcNAc by reverse hydrolysis activity of CmNAGase. GlcNAc of 10 g/L was incubated with CmNAGase (3 μ g) in 1 mL volume at pH 5.4 and 35°C for different time intervals. 1-6: Standards of GlcNAc, (GlcNAc)₂, (GlcNAc)₃, (GlcNAc)₄, (GlcNAc)₅, and (GlcNAc)₆.

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