A unique GCN5-related glucosamine N-acetyltransferase region exist in the fungal multi-domain glycoside hydrolase family 3 β-N-acetylglucosaminidase

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Glycoside hydrolase (GH) family 3 β-N-acetylglucosaminidases widely exist in the filamentous fungi, which may play a key role in chitin metabolism of fungi. A multi-domain GH family 3 β-N-acetylglucosaminidase from Rhizomucor miehei (RmNag), exhibiting a potential N-acetyltransferase region, has been recently reported to show great potential in industrial applications. In this study, the crystal structure of RmNag was determined at 2.80 Å resolution. The three-dimensional structure of RmNag showed four distinctive domains, which belong to two distinguishable functional regions — a GH family 3 β-N-acetylglucosaminidase region (N-terminal) and a N-acetyltransferase region (C-terminal). From structural and functional analysis, the C-terminal region of RmNag was identified as a unique tandem array linking general control non-derepressible 5 (GCN5)-related N-acetyltransferase (GNAT), which displayed glucosamine N-acetyltransferase activity. Structural analysis of this glucosamine N-acetyltransferase region revealed that a unique glucosamine binding pocket is located in the pantetheine arm binding terminal region of the conserved CoA binding pocket, which is different from all known GNAT members. This is the first structural report of a glucosamine N-acetyltransferase, which provides novel structural information about substrate specificity of GNATs. The structural and functional features of this multi-domain β-N-acetylglucosaminidase could be useful in studying the catalytic mechanism of GH family 3 proteins.

The general control non-derepressible 5 (GCN5)-related N-acetyltransferases (GNATs) are widespread in all kingdoms of living organisms and perform several important cellular functions through the acetylation of small molecules and protein substrates1. GNATs catalyse the transfer of acetyl groups from acetyl coenzyme A (AcCoA) to primary amine acceptors including aminoglycoside antibiotics2,3, glucosamine-6-phosphate (GlcN-6P)4,5, histones6,7, peptid8, arylamine9,9 and spermidine10. They were first found as aminoglycoside N-acetyltransferases in bacteria that develop antibiotic resistance to kanamycin and gentamicin1. Generally, there is very limited sequence conservation among members of the GNAT superfamily, in part, reflecting their capacity to bind and modify a diverse array of substrates. Previous structural studies revealed that although different enzymes of this superfamily show only moderate pairwise sequence homology, they share a common core fold comprising a central highly curved mixed β-sheet flanked on both sides by α-helices1-2.

Glucosamine N-acetyltransferase (EC 2.3.1.3) catalyses the transfer of an acetyl group from AcCoA to a glucosamine (GlcN) acceptor generating a N-acetylglucosamine. It has firstly been identified in pigeon liver extracts11.

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and participated in aminosugar metabolism. To date, these enzymes also have been found in plants and bacteria. The glycan chains of the peptidoglycan of the bacterial cell wall are composed of alternating β-1,4-glycosidically linked amino sugars N-acetylglucosamine (GlcNAc) and N-acetyl-muramic acid (MurNAc). Glucosamine N-acetyltransferases from some bacteria can recover non-N-acetylated sugars (the N-acetylation of GlcN residues at the non-reducing end of peptidoglycan fragments) prior to β-N-acetylglucosaminidase cleavage. However, there is no analogous glucosamine N-acetyltransferase structure that has been reported previously. In addition, whether the structural basis for substrate-specific acetylation by the fungal glucosamine N-acetyltransferase differs from the other GNATs is not clear.

The full-length β-N-acetylglucosaminidase (RmNag) from Rhizomucor miehei belongs to glycoside hydrolase (GH) family 3 in the CAZy classification of carbohydrate-active enzymes and the characterisation of which has been reported in our previous study. RmNag is a 858-amino-acid protein with multiple distinctive structural domains and the β-N-acetylglucosaminidase activity lies in the N-terminal region. Several structures of bacterial homologues of the glycoside hydrolase domains, involved in bacterial cell wall turnover, spore germination, and induction of β-lactamase, have been determined. These structures have been beneficial to design and analyse the inhibitors of GH family 3 β-N-acetylglucosaminidases. For the C-terminal region of RmNag, there is no information available at present although it may constitute an acetyltransferase region. In the present study, the crystal structure of full-length RmNag has been resolved to 2.8 Å resolution to find that a unique GCN5-related N-acetyltransferase region exists in the C-terminal region of RmNag, which displays glucosamine N-acetyltransferase activity in vitro. Our results not only present new information on the GCN5-related N-acetyltransferase family, but also provide a new assembly of a multi-domain β-N-acetylglucosaminidase. This is the first report on a crystal structure of a fungal GH family 3 β-N-acetylglucosaminidase, representing a significant advance in our understanding of these enzymes.

Results and Discussion

Overall structure. The crystal structure of full-length RmNag was determined to 2.80 Å resolution. The crystallographic statistics for data collection and structure refinement are given in Table 1. The monoclinic space group of RmNag was P42_21 with two monomers in the asymmetric unit. The overall structure of RmNag is presented in Fig. 1. The RmNag monomer with approximate dimensions of 90.7 × 50.1 × 77.5 Å, consisted of a series of four separate domains (A, B, C and D). The crystal structure of full-length RmNag reveals two distinct functional regions: a β-N-acetylglucosaminidase N-terminal region (NTM, domains A and B, residues 26–529), and a N-acetyltransferase C-terminal region (CTR, domains C and D, residues 537–858) (Fig. 1). The native molecular mass of RmNag as estimated by size-exclusion chromatography (SEC) is approx. 230 kDa (data not shown), suggesting that RmNag exists as a dimer in solution. However, the β-N-acetylglucosaminidase region (NTM) and the N-acetyltransferase region (CTR) are monomeric and dimeric in solution, respectively, indicating that full-length RmNag dimerizes via the interactions owing to its CTR. Indeed, a conserved dimer interface was observed in both the crystal structures of the full-length RmNag (Fig. 1) and its CTR. Even though the buried surface area in the interface of the full-length RmNag is relatively small (900 Å2), it harbors 16 hydrogen bonds and 8 salt bridges. It is noteworthy that total three sulfate ions were found in the crevice of domain D of monomers, and a binding AcCoA molecule was found in the each crevice of domain C of monomers.

β-N-Acetylglucosaminidase N-terminal region (NTM). The β-N-acetylglucosaminidase N-terminal region (NTM) of RmNag covers two domains (A and B) (Fig. 2a). The N-terminal domain (domain A, residues 26–371) reveals a characteristic (β/α)8 TIM barrel, which is typical for catalytic domains of GH family 3 members (Fig. 2a). The C-terminal domain (domain B, residues 372–529) displays a 3α/6β/3α sandwich-like fold, in which a six-stranded β-sheet is sandwiched by two layers of three helices each (Fig. 2a). Even though domain B is tightly associated with domain A, domain B is located fairly far away from the catalytic site of domain A. Hence, the domain B’s function remains unclear but it is unlikely to be directly involved in the β-N-acetylglucosaminidase activity. An approximately 9.2 Å long and 5.5 Å deep catalytic pocket is located on the reverse side of the NTM molecule, allowing the binding of N-acetylglucosaminic acid. The center of this catalytic pocket is surrounded by the glyceraldehyde 3-phosphate dehydrogenase A. In several structures of multi-domain GH family 3 β-N-acetylglucosaminidases, the conserved aspartate residue from the (β/α)8 TIM barrel domain and a general acid/base catalytic glutamate residue from the 3α/6β/3α sandwich domain have been proposed to function as the catalytic acid/base, which is unique for GH family 3 members. In a recent study suggests that these enzymes may act preferentially as glycoside phosphorylases. Histidine is used as acid-base catalyst in place of the aspartate residue identified in several structures of multi-domain GH family 3 β-N-acetylglucosaminidases. Thus, it cannot be assumed that all GH family 3 β-N-acetylglucosaminidases are glycoside phosphorylases at present.

In the structure of RmNag, a non-canonical His192/Asp190 dyad able to function as the catalytic acid/base is present on a flexible loop which is 6.3 Å apart from the conserved aspartate nucleophile (Asp268) (Fig. 2a). Furthermore, the N1 of His192 forms a hydrogen bond with Oε of Asp190 at a distance of 2.9 Å. A similar arrangement of the Asp-His-Asp triad has been reported for the β-N-acetylglucosaminidase of Bacillus subtilis. Superposition of the β-N-acetylglucosaminidase region (NTM) onto the Bacillus subtilis β-N-acetylglucosaminidase...
in complex with PUGNAc (O-(2-acetamido-2-deoxy-D-glucopyranosylidene) amino N-phenylcarbamate; PDB accession code 3NVD) indicates a conserved overall fold and catalytic site (Fig. 2b). The overall root-mean-square deviation (RMSD) value for 513 superimposed C$\alpha$ atom pairs is 1.3 Å. The importance of catalytic residues His192 and Asp268 was corroborated by site-directed mutagenesis, in which substitution of each of these residues for alanine (H192A or D268A) clearly prohibited the enzyme’s hydrolysis activity (Table 2). The structural study of RmNag provides another example that atypical subfamily $\beta$-N-acetylglucosaminidases within the GH family 3 members might use a His/Asp dyad as the catalytic acid/base.

### N-Acetyltransferase C-terminal region (CTR).

The N-acetyltransferase C-terminal region (CTR) of RmNag contains two structurally conserved GNAT-fold domains (C and D). A N-terminal $\beta$ strand is followed by two $\alpha$ helices, three antiparallel $\beta$ strands, and then followed by a signature central $\alpha$ helix, a fifth $\beta$ strand, a fourth

### Table 1. Data collection and refinement statistics.

Values in parentheses represent the highest resolution shell. $^a R_{\text{merge}} = \sum_{hkl} \sum_i |I(hkl)_i - \langle I(hkl) \rangle| / \sum_{hkl} \sum_i I(hkl)_i$, where $I(hkl)_i$ is the intensity of reflection $hkl$ and its symmetry equivalents and $\langle I(hkl) \rangle$ is the average intensity over all equivalent reflections. $^b R_{\text{measure}}$: multiplicity-weighted $R_{\text{merge}}$; $R_{\text{free}}$: precision-indicating $R_{\text{merge}}$. $^c R = \sum |F_o| - |F_c| / \sum |F_o|$. $|F_o|$ and $|F_c|$ are amplitudes of the observed and calculated structure factors, respectively. $R_{\text{work}}$ is the $R$ value for reflections used in refinement, whereas $R_{\text{free}}$ is the $R$ value for 5% of the reflections which are selected randomly and are not included in the refinement.

| Data collection statistics | CTR, Se-derivative | RmNag, full-length |
|---------------------------|---------------------|------------------|
| X-ray source              | BL14.2, BESSY, Berlin | P11, PETRA III, Hamburg |
| Detector                  | MX-225 CCD          | Pilatus 6M fast   |
| Wavelength (Å)            | 0.9798              | 1.0331           |
| Space group               | $P4_22$             | $P4_22$          |
| Unit-cell dimensions (Å)  | $a = b = 82.16, c = 180.03$ | $a = b = 245.03, c = 94.52$ |
| Resolution range (Å)     | 49.05–2.26 (2.38–2.26) | 88.19–2.80 (2.95–2.80) |
| Number of unique reflections | 30364 (4277)         | 71082 (10201)    |
| Completeness (%)          | 99.8 (98.5)         | 99.9 (99.3)      |
| Mean I/sigma(I)           | 19.1 (2.6)          | 29.4 (4.3)       |
| Multiplicity              | 26.9 (27.0)         | 26.2 (27.3)      |
| $R_{\text{merge}}(\%)^a$ | 11.4 (140.9)        | 11.8 (78.6)      |
| $R_{\text{free}}(\%)^b$  | 11.6 (143.6)        | 12.0 (80.0)      |
| $R_{\text{free}}(\%)^c$  | 2.2 (27.3)          | 2.3 (15.1)       |
| Wilson B-factor (Å$^2$)   | 40.2                | 62.6             |

| Refinement statistics     | CTR, Se-derivative   | RmNag, full-length |
|---------------------------|----------------------|-------------------|
| Resolution range (Å)      | 86.63–2.80           |                   |
| No. of reflections in working/test | 66830/3561           |                   |
| $R_{\text{work}}(\%)^c$  | 22.9 (28.9)          |                   |
| $R_{\text{free}}(\%)^c$  | 25.4 (31.7)          |                   |
| Protein atoms             | 13603                |                   |
| No. of water molecules    | 114                  |                   |
| No. of ions               | 3                    |                   |
| RMSD in bond lengths (Å)  | 0.013                |                   |
| RMSD in bond angles (°)   | 1.70                 |                   |
| RMSD B-factor for bonded atoms | 2.67                 |                   |
| Average B-factor (Å$^2$)  | Overall 51.11        |                   |
|                           | Protein 50.55         |                   |
|                           | Solvent 54.16         |                   |
|                           | Ions 111.14           |                   |
|                           | Ligands 69.84         |                   |
| Ramachandran plot (%)     | Most favored 94.3     |                   |
|                           | Additionally allowed 5.2 |                   |
|                           | Outliers 0.5          |                   |
|                           | Rotamer outliers (%)  5.1 |                   |
|                           | C-beta outliers (%)   0.25 |                 |
|                           | Clashscore 6.76       |                   |
|                           | Overall score 2.29    |                   |

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α helix and a final β strand (Fig. 3a). In spite of less than 10% sequence identity, these two domains share an overall RMSD value of 2.6 Å (for 130 superimposed Cα atom pairs) (Figs 3c and 4c). It is noteworthy that the two analogous GNAT-fold domains are arranged in a tandem array linked to the C-terminal of the β-N-acetylglucosaminidase region. Domain C and domain D are linked by one α helix. Therefore, the N-acetyltransferase region (CTR) represents a member of the subfamily of tandem GNAT structures (Fig. 3a).

A co-purified AcCoA molecule was found in the N-terminal of CTR (domain C, Ala537–Leu693). This AcCoA tightly interacts with the domain C via extensive hydrogen bonds and some hydrophobic interactions (Fig. 3b). However, this AcCoA molecule and the domain C are unlikely to be directly involved in acetyl transfer, mainly because there are no residues nearby the acetyl group that can be possibly involved in general acid/base catalysis. Several mutagenesis data have proposed a tyrosine as the proton donor for the thiolate leaving group1,23. However, sequence alignments of other GNAT homologues with the domain C indicate that the conserved tyrosine is replaced by a phenylalanine in the corresponding position of domain C (Figs 3c and 4c). In addition, this AcCoA molecule was tightly bound and its acetyl group was found to be deeply buried. It is noteworthy that a loop 654PRFFPGVPDDAQ666 which links the fifth β strand and the fourth α helix of domain C is missing in other GNAT sequences (Fig. 3a,c). The insert loop may block the entrance of the active site of domain C and thus hinder the substrate to productively bind to the catalytic pocket (Fig. 4a).

Consequently, the acetyl transfer activity must be located within the C-terminal of CTR (domain D, Ile711–Phe858). A structural homology search at the DALI server revealed that RmNag domain D is similar in structure to Mycobacterium tuberculosis mycothiol synthase (PDB accession code 1OZP)24 with an RMSD of 2.4 Å (Supplementary Table S1), despite an extremely low amino-acid sequence identity (~16%). Relevant structures are the N-acetyltransferase from Sphaerobacter thermophilus (3TT2), the N-acetyltransferase from Trypanosoma brucei (3FB3), the putative acetyltransferase from Shigella flexneri (2PDO), the putative N-acetyltransferase from Escherichia coli (4QVT), the GlcN-6P N-acetyltransferase from Saccharomyces cerevisiae (1ID), the acetyltransferase from Listeria monocytogenes (2OH1), and the hypothetical protein from Drosophila melanogaster (1SQH). The structure of GNATs is conserved from ancestral to prokaryotic and eukaryotic cells, although they show different
types of substrate specificity. Superposition of the HsGNA1–CoA–GlcNAc-6P complex (2OZ8) onto the domains C and D showed that domain D is structurally more conserved than domain C, with overall RMSD values for 130 Cα atom pairs of 1.3 Å and 2.3 Å, respectively (Fig. 4c). The substrate-binding pocket of domain D is clearly divided into two parts: the positive charged and negative charged clefts are docked with CoA and GlcNAc molecules, respectively (Fig. 4b). Most important, the conserved tyrosine (Y842 in RmNag) which has been predicted to act as the general acid to protonate the leaving thiolate anion nearly overlapped with the one of the HsGNA1–CoA–GlcNAc-6P complex and it is therefore within hydrogen bond distance from the sulphur atom of a bound CoA (Fig. 4c). Mutation studies show that RmNag Y842A lost its transacetylation activity (Table 2), which confirmed the inference above.

![Figure 2. Structure of the β-N-acetylglucosaminidase region (NTR).](image)

(a) Cartoon representation of the β-N-acetylglucosaminidase region (NTR). The α-helices, β-strands, and loops of domain A of RmNag are coloured in red, yellow, and green, respectively and those of domain B of RmNag are coloured in cyan, magenta, and orange, respectively. The catalytic residues (His192, Asp190, and Asp268) are presented as coloured sticks. (b) Superimposition of BsNag-PUGNAc complex (cyan, PDB: 3NVD) and RmNag β-N-acetylglucosaminidase region (magenta, NTR). The PUGNAc is shown in stick representation. Details of the active site geometry of NTR and BsNag-PUGNAc in comparison are shown in an amplified and detailed view. Active-site residues are shown as stick models. The residues of the NTR are shown in magenta and labelled. The residues of BsNag-PUGNAc are shown in cyan and labelled in brackets. (c) Electrostatic-potential surface representation of the β-N-acetylglucosaminidase region (NTR) of RmNag (upper half). Red and blue surfaces represent negative and positive potentials (–10 kBT to +10 kBT), respectively. The electrostatic surface potentials were calculated using the Adaptive Poisson-Boltzmann Solver (APBS)47 within the APBS plugin for PyMOL (v.1.3; Schrödinger LLC).

| RmNag     | Specific activity (U mg⁻¹) | Relative activity (%) |
|-----------|----------------------------|-----------------------|
| Wild type | Hydrolysis activity        |                       |
|           | 29.0                       | 100                   |
| NTR       | 22.7                       | 78.3                  |
| CTR       | None*                      | 0                     |
| D268A     | None                       | 0                     |
| H192A     | None                       | 0                     |
|           | Transacetylation activity  |                       |
| Wild type | 115.2                      | 100                   |
| CTR       | 57.7                       | 50.1                  |
| NTR       | None                       | 0                     |
| Y842A     | None                       | 0                     |

Table 2. Hydrolysis and transacetylation activities of wild type and variants of RmNag. *No activity was detected.
In total, the structural analysis suggests that the AcCoA in the domain C might only possess a structural function, whereas the domain D is likely directly involved in acetyl transfer.

N-Acetyltransferase properties of \textit{RmNag}. The N-acetyltransferase properties of full-length \textit{RmNag}, NTR and CTR toward GlcN as a substrate were investigated by thin-layer chromatography (TLC). The full-length \textit{RmNag} and CTR could transfer the acetyl to GlcN using AcCoA as the co-substrate, releasing GlcNAc as final product (Fig. 5a). However, the NTR had no N-acetyltransferase activity since GlcN and AcCoA did not react in the presence of NTR. N-acetyltransferase activity of the full-length \textit{RmNag}, NTR and CTR were then assayed by high performance liquid chromatography (HPLC). Full-length \textit{RmNag} and CTR exhibited N-acetyltransferase activity of 115.2 and 57.7 U mg$^{-1}$, respectively. In contrast, no N-acetyltransferase activity was detected by NTR (Table 2). On the other hand, full-length \textit{RmNag} and NTR exhibited \(\beta\)-N-acetylglucosaminidase activity of 29.0 and 22.7 U mg$^{-1}$, respectively. And no \(\beta\)-N-acetylglucosaminidase activity was detected by CTR. This result confirmed that the CTR is the N-acetyltransferase region of \textit{RmNag}. Furthermore, \textit{RmNag} had N-acetyltransferase activity towards both GlcN and chitosan-oligomers. It exhibited N-acetyltransferase activity range of 115.2 to 5.4 U mg$^{-1}$ towards GlcN to (GlcN)$_5$ (Table 3).

To detect the unique catalytic properties of this GH family 3 enzyme, the hydrolysis properties of full-length \textit{RmNag} towards (GlcN)$_2$ were further investigated (Fig. 5b). TLC analysis showed that the substrate (GlcN)$_2$ declined gradually in the process of the enzymatic reaction, when AcCoA was present in the reaction mixture. Meanwhile, the products (GlcNAc)$_2$ and GlcNAc increased gradually. Full-length \textit{RmNag} almost completely acetylated and hydrolyzed all of the newly produced (GlcN)$_2$ in 4 h to yield the end product. These results indicated that full-length \textit{RmNag} could transfer the acetyl to (GlcN)$_2$ using AcCoA as the co-substrate to produce (GlcNAc)$_2$. 

Figure 3. Structure of the N-acetyltransferase region (CTR). (a) The overall structure of the acetyltransferase region (CTR) of \textit{RmNag}. Domains C and D are coloured in red and cyan, respectively. The linker \(\alpha\)-helix is coloured in yellow. The insert loop 654PRFFPGVPDDAQ666 of domain C (absent in domain D) is highlighted in magenta. (b) The AcCoA ligand interacts extensively with domain C. Residues involved in the interactions are shown as coloured sticks (AcCoA carbon atoms: white; domain C carbon atoms: green; N: blue; O: red; P: orange; S: yellow) and labelled. Hydrogen bonds are represented by red dashed lines. (c) Sequence alignment of domains C and D with other GNAT homologues. The alignments were done using MUSCLE$^{45}$ and the coloured figures were generated using ESPript3.0$^{46}$. Residues coloured in red are conserved to more than 70% and residues boxed in red are completely identical. Secondary structures of domains C and D are indicated on top and bottom of the alignments, respectively. The tyrosine that is considered as a putative general acid is indicated by a star symbol. All the sequences aligned here were taken from the Genbank database (http://www.ncbi.nlm.nih.gov/genbank/): \textit{Saccharomyces cerevisiae} GANT (ScGNAT, GenBank ID: NP_116637.1), \textit{Aspergillus fumigatus} GNAT (AfGNAT, GenBank ID: XP_747831.1), \textit{Homo sapiens} GNAT (HsGNAT, GenBank ID: NP_932332.1), \textit{Mus musculus} (MmGNAT, GenBank ID: NP_062298.1).
Figure 4. **Structural comparison of the two acetyltransferase folds.** (a) Electrostatic-potential surface of domain C of *RmNag*. The substrate-binding pocket of domain C is occupied by a co-purified AcCoA. Red and blue surfaces represent negative and positive potentials (–10 kBT to + 10 kBT), respectively. The electrostatic surface potentials were calculated using the APBS within the APBS plugin for PyMOL (v.1.3; Schrödinger LLC). A 90° rotation view of the substrate-binding pocket indicates that the insert loop may block the entrance of activity site of domain C. (b) Electrostatic-potential surface of domain D of *RmNag*. CoA and GlcNAc molecules are docked into the substrate-binding pocket of domain D. (c) Superimposition of domains C (red) and D (cyan) (left); domain C (red) and *HsGNA1*-CoA-GlcNAc-6P complex (green; PDB: 2O28) (middle); domain D (cyan) and *HsGNA1*-CoA-GlcNAc-6P complex (green) (right). The AcCoA, CoA and GlcNAc-6P ligands are shown as stick models (AcCoA from domain C: C: yellow, N: blue, O: red, S: brown; CoA and GlcNAc-6P from 2O28: C: white, N: blue, O: red, P: orange, S: brown). The difference of orientation between the thioester group of AcCoA from domain C and the thiol group of CoA from *HsGNA1*-CoA-GlcNAc-6P complex (2O28) are shown in the magnified view. The insert loop of domain C is highlighted in magenta.

Figure 5. **TLC analyses of the transacetylation and hydrolysis products by *RmNag*.** (a) The transacetylation properties of *RmNag* (full length, NTR and CTR). Lane 1: marker, GlcNAc; lane 2: marker, GlcN; lane 3: control, reaction mixture without AcCoA; lane 4: control, reaction mixture without enzyme; lane 5: reaction mixture, full length *RmNag*; lane 6: reaction mixture, CTR; lane 7: reaction mixture, NTR. (b) Reaction process of chitobiose ((GlcN)_2) hydrolysed by *RmNag* (full length). M: marker sugars. During the reaction progress, (GlcN)_2 was tapered off and GlcNAc and (GlcNAc)_2 were gradually accumulated.
The N-acetyltransferase product was then cleaved by full-length *Rm* Nag at the β-1,4 linkage in the meantime, to further yield GlcNAc monomer as the final product.

In our previous study, *Rm* Nag cleaved (GlcNAC)_{n}, in the absence of AcCoA. However, *Rm* Nag could not cleave (GlcN)_{n} in the absence of AcCoA (Supplementary Fig. S1). When AcCoA was present in the reaction mixture, *Rm* Nag cleaved (GlcN)_{n} to produce GlcNAc (Fig. 5b). Furthermore, both full-length and NTR displayed N-acetylglucosaminidase activity towards pNP-GlcNAc, but the CTR had no N-acetylglucosaminidase activity (Table 2). Thus, the transferase activity is not required for the N-acetylglucosaminidase to cleave (GlcNAc)_{n}, but is processive and required for the N-acetylglucosaminidase to cleave (GlcNAc)_{n}.

**N-Acetyltransferase properties**

The transferase properties were in accord with that of the glucosamine N-acetyltransferase from *Clostridium acetobutylicum* [13]. It is interesting that owing to the unique CTR, *Rm* Nag could hydrolyse chitobiose to produce acetylglucosaminide. This property is similar with the exo-β-D-glucosaminidase (EC 3.2.1.165) involved in chitosan hydrolysis [25,26]. Previous studies showed that some O-GlcNAc hydrolizing enzymes (O-GlcNAcases) are also multi-domain proteins [27,28], which exhibit glycoside hydrolase activity in the N-terminal domain and have a C-terminal domain with low sequence similarity to known acetyltransferases. The N-terminal region of O-GlcNAcases is a GH family 84 β-N-acetylglucosaminidase domain which specifically catalyzes the cleavage of GlcNAc from modified proteins. The C-terminal region of O-GlcNAcases is a GNAT domain which may function as a histone acetyltransferase (albeit controversial) [27,28]. To date, there is no information available on how the domains of these multi-domain β-N-acetylglucosaminidases interact and how the substrate is delivered from one to the other active site. The active sites of the NTR and CTR of *Rm* Nag are separated quite clearly. Also, there is no interaction of functional sites at the interfaces of the different domains derived from the current structural information of *Rm* Nag. It is thus speculated that the different domains of *Rm* Nag only possess a structural connection. The acetylated product is released into the reaction system by the N-acetyltransferase region, and then accesses the active site of the β-N-acetylglucosaminidase domain for hydrolytic cleavage. To probe this hypothesis, effects of pH and temperature on the glucosamine N-acetyltransferase activity of *Rm* Nag were examined. *Rm* Nag displayed maximal glucosamine N-acetyltransferase activity at pH 6.5 and exhibited optimal activity at 55 °C (Supplementary Fig. S2). This corresponds to the optimal pH and temperature of β-N-acetylglucosaminidase activity of *Rm* Nag, which are pH 6.5 and 50 °C, respectively [16]. These results illustrate that the β-N-acetylglucosaminidase and glucosamine N-acetyltransferase activities of *Rm* Nag are carried out at similar environmental conditions.

**Structural basis of substrate specificity of CTR.**

The GNAT superfamily is one of the largest enzyme superfamilies recognized to date and has more than 10,000 representatives from all kingdoms of life [1]. In spite of modest degrees of overall primary sequence homology, the basic structure of the GNAT fold is extraordinarily conserved, and serves two nearly universal functions: to bind the pantetheine arm of AcCoA and to polarize the carbonyl of the thioester through hydrogen bond interactions. However, diverse GNAT members have different substrate specificities, which play important roles in life activities. Therefore it is imperative to identify structural differences among various GNATs. There is no structural information available on enzymes that are functionally homologous to glucosamine N-acetyltransferases. The crystal structure of *Rm* Nag allowed us to address the molecular details of substrate binding and catalysis of the glucosamine N-acetyltransferase being different from other GNAT superfamily members. A superposition of *Rm* Nag domain D with typical GNAT complexes showed that the majority of GNATs have similar CoA binding pockets (Fig. 6a). However, the acceptor binding pockets were totally different among various GNATs. These different regions confer the substrate specificities of various GNATs. The superposition result showed that GlcN-6P N-acetyltransferase, spermidine/spermine N1-acetyltransferase and serotonin N-acetyltransferase have a loop region near the pantetheine arm terminal of CoA, which provides the residues for substrate binding (Fig. 6a). In contrast, domain D of *Rm* Nag lacked this substrate-binding loop. In addition, an α-helix (α2) existing in this region implied that the N-acetyltransferase domain of *Rm* Nag would show a unique substrate binding mode among GNATs. To reveal the structural basis of substrate specificity of CTR, the superposition of CTR with the GlcN-6P N-acetyltransferase (2O28) is shown in Fig. 6b. GlcN-6P N-acetyltransferase catalyses the transfer of acetyl groups from AcCoA and recognizes analogous acceptors compared to the glucosamine N-acetyltransferase (GlcN and GlcN-6P, respectively). The substrate binding pocket of the GlcN-6P N-acetyltransferase can be divided into two parts: a GlcNAc binding pocket (Lys108, Asp121, Val122 and Glu156) and a phosphate binding pocket (Thr61, His111, Tyr151, Lys152 and Arg181). In contrast, CTR lacks the entire phosphate-binding pocket (Fig. 6b). This structural architecture excludes CTR from binding GlcN-6P. On the other side, CTR possesses a GlcNAc binding pocket, which is similar to the GlcN-6P N-acetyltransferase.

### Table 3. Transacetylation activity of *Rm* Nag towards GlcN and chitosan-oligomers.

| Substrate | Specific activity (U mg⁻¹) | Relative activity (%) |
|-----------|---------------------------|----------------------|
| GlcN      | 115.2                     | 100                  |
| (GlcN)₅   | 89.3                      | 77.5                 |
| (GlcN)₆   | 13.8                      | 12.0                 |
| (GlcN)₇   | 10.7                      | 9.3                  |
| (GlcN)₈   | 5.4                       | 4.7                  |

*Specific activity towards chitosan-oligomers was determined by HPLC. Reactions were performed in 50 mM Bis-Tris buffer pH 6.5 at 55 °C for 10 min.*
Residues Thr738, Cys799, Val800 and Asp834 of CTR superpose on the corresponding residues of the GlcN-6P N-acetyltransferase’s GlcNAc binding pocket. Furthermore, two aromatic residues (Trp735 and Trp835) are located in the GlcNAc binding pocket of CTR, which may stack against the pyranose ring, forming the hydrophobic sugar-binding platform (Fig. 6b). This architecture forms the structural basis of substrate specificity of CTR to bind glucosamine.

CTR consists of two fused GNAT domains, only one of which is functional. This cross-talk of the domains is less reported in GNATs. The other two reported tandem GNATs are Saccharomyces cerevisiae N-myristoyltransferase\textsuperscript{30} and Mycobacterium tuberculosis mycothiol synthase\textsuperscript{24}. On the basis of the two previous studies and the structural information of Rm Nag, the function of the AcCoA ligand bound to the N-terminal domain (domain C) is still unclear; it may act as an effector molecule or function to stabilize the domain\textsuperscript{24,30}. Furthermore, some tandem GNATs (2O28 and 1OZP) form unique active sites at their dimer interfaces. The similarity of the subunit interface of typical GNAT proteins and the interface between the two GNAT domains of these tandem GNATs suggests that the progenitor of tandem GNAT may have arisen from gene duplication and fusion of a homodimeric GNAT, followed by structural rewiring through mutation and selection\textsuperscript{24}.

Phylogenetic and sequence analyses of Rm Nag. Typical GH family 3 members consist of two diverse domains\textsuperscript{31,32}. The GH family 3 β-N-acetylglucosaminidases have been proved to possess an Asp–His–Asp triplet catalytic core\textsuperscript{17,21}. Unlike other members of the GH family 3, all of the three catalytic residues were detected in the N-domain of the enzymes (β/α)\textsubscript{8} TIM barrel domain). Some β-N-acetylglucosaminidases of the GH family 3 even completely lack a C-domain\textsuperscript{33}. Phylogenetic analysis of GH family 3 β-N-acetylglucosaminidase (or putative β-N-acetylglucosaminidase) ies (Supplementary Fig. S3). Sequences generated a tree in which all 41 sequences were placed into two subfamilies. Subfamily 1 contained 21 sequences, including several β-N-acetylglucosaminidases from gram negative bacteria, such as Salmonella typhimurium, Escherichia coli, and Burkholderia cenocepacia. It is noteworthy that all three structurally characterized β-N-acetylglucosaminidases of subfamily 1 are single domain β-N-acetylglucosaminidases. All the members of subfamily 1 lack the C-terminal domain sequences which are typical of GH family 3 proteins. Subfamily 2 covered 20 sequences, including Rm Nag and other proteins from fungi, gram positive bacteria and microalgae. Subfamily 2 members are multi-domain proteins, which contain C-terminal domain sequences of other representative GH family 3 proteins. However, different from other typical GH family 3 proteins, the C-terminal domains of subfamily 2 β-N-acetylglucosaminidases lacked the catalytic residues and thus did not take part in the catalysis. The subfamilies can be distinguished by differences in the sequence pattern next to the conserved P/V(V)L/D motif in the N-terminal domain. Rm NAG and other β-N-acetylglucosaminidases from fungi possess an additional region that has been predicted to be a N-acetyltransferase region. These fungal proteins can be divided into an independent clade from subfamily 2. This unique glucosamine N-acetyltransferase region was found to widely exist in the C-terminal of GH family 3 β-N-acetylglucosaminidases from fungi such as Mucor, Rhizopus, Aspergillus, Penicillium, and Parastitella deposited at the Genebank.
Physiological role of RmNag. From the N-acetyltransferase properties and sequence analyses of RmNag, the most possible physiological function of RmNag is an involvement in the fungal chitin metabolism and cell wall rearrangement. Chitin, composed of β-1,4-linked N-acetylglucosamine (GlcNAc) units, is present as an important component in the cell wall of fungi, which is essential for fungi to maintain cell structure integrity. The acetylation level of the fungal cell wall chitin is dynamic and the chitin polymer usually contains some GlcN units. In the fungal chitin metabolism, terminal GlcN units from chitin (or related oligosaccharides) are hardley cleaved by β-N-acetylglucosaminidases. Therefore, an exo-β-D-glucosaminidase (EC 3.2.1.165) activity is needed in the fungal chitin metabolism.

According to the classification of carbohydrate-active enzymes (CAZY), exo-β-D-glucosaminidases are grouped into three GH families: 2, 9, and 35. However, no fungal exo-β-D-glucosaminidase has been characterized to date. On the basis of the above consideration, multi-domain β-N-acetylglucosaminidases may play this role in fungal chitin metabolism instead of an exo-β-D-glucosaminidase. This is consistent with findings that the unique glucosamine N-acetyltransferase region widely exists in the C-terminal region of GH family 3 β-N-acetylglucosaminidases from fungi.

Conclusions
In this study, the full-length multi-domain GH family 3 β-N-acetylglucosaminidase from R. miehei (RmNag) was structurally characterized to resolution of 2.80 Å. The crystal structure of RmNag displays four separate domains, which belong to two distinct functional regions. From the three-dimensional structure of RmNag, a unique tandem array linking GCN5-related N-acetyltransferase region exists in the C-terminal of RmNag. This N-acetyltransferase region was identified as a glucosamine N-acetyltransferase region by structural and functional analysis. The structural basis for its substrate binding pocket, which is feature required for the proper function of this protein, has been identified based on structural homology within the GNAT members. These results should be useful in studying the catalytic mechanism of other GH family 3 proteins and provide novel information on the GCN5-related N-acetyltransferases.

Methods
Cloning, gene expression and protein purification. DNA sequences encoding full-length RmNag, N-terminal region (NTR, Met1–Asn529) and C-terminal region (CTR, Val530–Phe858), were amplified by PCR using Pfu DNA polymerase (Invitrogen) from the genomic DNA of R. miehei CAU432. The resulting fragments were then cloned into a modified Sumo-pET28a(+) expression vector using BamHI and NotI sites. All genes were expressed in the Escherichia coli strain BL21-Gold (DE3). Briefly, the cultures were grown in 2×YT medium at 37°C until the optical density at 600 nm was up to 0.8. Expression was induced by adding isopropyl-β-D-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM and incubation at 25°C overnight. The selenomethionyl (SeMet) derivative of CTR was prepared using the method of methionine-biosynthesis pathway inhibition. Cells were harvested by centrifugation and lysed by sonication in buffer A (50 mM Tris–HCl pH 8.0, 20 mM imidazole and 300 mM NaCl). The lysate was ultracentrifuged at 20,000 rpm for 60 min at 4°C, and the supernatant was applied onto a pre-equilibrated 5 mL HisTrap HP column (GE Healthcare), followed by washing with 100 mL buffer A. The proteins were eluted with buffer B (50 mM Tris–HCl pH 8.0, 300 mM NaCl and 500 mM imidazole), and the eluted samples were mixed with SUMO-protease. The cleavage was performed at 4°C overnight. The cleaved samples were further purified by size-exclusion chromatography (HiLoad 16/60 Superdex 200; GE Healthcare) equilibrated with buffer C (10 mM Tris–HCl pH 7.5, 150 mM NaCl and 5 mM DTT). The recombinant proteins contain two additional amino acids (Gly and Ser) at their N-terminals.

The site-directed mutagenesis (D268A, H192A and Y842A) were performed directly on the Sumo-pET28a(+) expression vector containing the RmNag gene, by the quick-change method with the Fast Mutagenesis System site-directed mutagenesis kit (TransGen Biotech, China). Primers used in gene amplification and site-directed mutagenesis are listed in Supplementary Table S2. The desired mutants were selected, sequenced and transformed into E. coli strain BL21-Gold (DE3) for expression.

Crystallization and data collection. Crystals were screened using a Phoenix robot (Art Robbins Instruments) with the sitting-drop, vapor-diffusion method at 20°C and drops containing 0.3 μL of protein solution plus 0.3 μL of reservoir solution. The following commercial screening kits were used: SaltRxTM, PEG/IonTM, IndexTM, Crystal ScreenTM, and PEGRxTM (all from Hampton Research). Crystals of both RmNag full-length and CTR appeared in a drop containing 0.1 M Bis-Tris (pH 5.5–6.5), 2 M (NH₄)₂SO₄. Optimized crystals suitable for diffraction were grown in drops containing 1.5 μL of protein solution and 1.5 μL of reservoir solution (1.3–1.6 M (NH₄)₂SO₄, 0.1 M Bis-Tris pH 5.5–6.5) at 20°C. Crystals were soaked in reservoir solution supplemented with 20% glycerol, and then vitrified in liquid nitrogen. Diffraction data of the CTR and full-length RmNag were collected at 100 K using beamline BL14.2 at BESSY (Berlin, Germany) and P11 at PETRA III (Hamburg, Germany), respectively. Indexing, integration and scaling of data were carried out with XDS. The program XPREP (Bruker) was used to further analyze and prepare datasets for structure solution and refinement. Statistics of the datasets are summarized in Table 1.

Phase determination, model building and refinement. The crystal structure of the RmNag CTR was determined by the single-wavelength anomalous diffraction (SAD) method. Phase calculations and initial model building were carried out by the program AutoSol from the PHENIX suite. Thereafter, model building and refinement were performed using Coot and Refmac5, respectively. To determine the RmNag full-length structure, a model representing its NTR was prepared firstly by modifying the coordinates of Bacillus subtilis N-acetylglucosaminidase (PDB code: 2BMX), guided by an amino acid sequence alignment. The resultant NTR model, together with the model of CTR, were employed as search models in PHASER. The refinement was
Enzyme assay and transacetylation properties. N-acetyltransferase activity was determined by high performance liquid chromatography (HPLC) using glucosamine as the substrate (acceptor) and N-acetylglucosamine as the product. The reaction mixtures containing 20 μL of 50 mM substrate in 50 mM Bis-Tris pH 6.5, 20 μL of 50 mM AcCoA (as acetyl donor) and 20 μL of suitably diluted enzyme (full length or CTR) were incubated at 55°C for 10 min. The reactions were then terminated by boiling for 5 min and substrate/product determined by HPLC (Agilent1200, Agilent, USA) equipped with a gel-filtration sugar–D KS-802 column (Shodex, Japan). The N-acetylglucosamine was eluted with water at a flow rate of 0.6 mL min⁻¹ at 65°C and the absorbance monitored at 210 nm. One unit of enzyme activity was defined as the amount of enzyme required to produce 1 μmol N-acetylglucosamine per minute under the above mentioned conditions.

β-N-acetylglucosaminidase activity was determined spectrophotometrically with pNP-GlcNAc as the substrate. One unit of enzyme activity was defined as the amount of enzyme required to liberate 1 μmol of pNP per minute under the assay conditions.

The transacetylation properties of \( R_m \)Nag were investigated by analysis of reaction products from the glucosamine and chitobiose. To test the N-acetyltransferase activity of \( R_m \)Nag (full length, NTR and CTR), 20 μL of suitably diluted enzyme was added to 50 mM glucosamine in 50 mM Bis-Tris pH 6.5 with 20 μL of 50 mM AcCoA, and then incubated at 30°C for 4 h. Samples withdrawn at different times were immediately boiled for 5 min, and then analysed by thin-layer chromatography (TLC). The control samples were reaction mixtures without AcCoA or without enzymes, and incubated at the same conditions. The enzymatic reaction supernatants were spotted onto TLC plates (silica gel 60 F₂₅₄; Merck, Germany) using n-butanol:methanol:ammonia:water (5:4:2:1 v/v/v/v) as the developing solution. The TLC plates were visualized after dripping the plate in methanol containing 2% (v/v) concentrated H₂SO₄, following by heating at 130°C for a few minutes. The synergy of N-acetyltransferase and β-N-acetylglucosaminidase activity of \( R_m \)Nag was investigated by analyzing the hydrolytic products from the chitobiose. Reaction mixtures (10 μL) containing 50 mM chitobiose and 50 mM AcCoA were incubated in 50 mM Bis-Tris pH 6.5 with suitably diluted enzyme (full length) at 30°C for 4 h. Samples withdrawn at different times were terminated by boiling for 5 min, then analysed by TLC as above.

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Author Contributions
Z.Q.J. and M.J.R. designed and supervised the project; Z.Q., Y.B.X., X.B.Y. and S.Q.Y. performed research; Z.Q.
and Y.B.X. wrote the manuscript. All the authors read and edited the manuscript, and approved the manuscript.

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Author Contributions
Z.Q.J. and M.J.R. designed and supervised the project; Z.Q., Y.B.X., X.B.Y. and S.Q.Y. performed research; Z.Q.
and Y.B.X. wrote the manuscript. All the authors read and edited the manuscript, and approved the manuscript.

Additional Information
Accession codes: The atomic coordinates and structure factors for the crystal structure of RnmNag have been
deposited at the Protein Data Bank (http://www.pdb.org) under accession code 4ZM6.

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