Synergy of Peptide and Sugar in O-GlcNAcase Substrate Recognition

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SUMMARY

Protein O-GlcNAcylation is an essential reversible posttranslational modification in higher eukaryotes. O-GlcNAc addition and removal is catalyzed by O-GlcNAc transferase and O-GlcNAcase, respectively. We report the molecular details of the interaction of a bacterial O-GlcNAcase homolog with three different synthetic glycopeptides derived from characterized O-GlcNAc sites in the human proteome. Strikingly, the peptides bind a conserved O-GlcNAcase substrate binding groove with similar orientation and conformation. In addition to extensive contacts with the sugar, O-GlcNAcase recognizes the peptide backbone through hydrophobic interactions and intramolecular hydrogen bonds, while avoiding interactions with the glycopeptide side chains. These findings elucidate the molecular basis of O-GlcNAcase substrate specificity, explaining how a single enzyme achieves cycling of the complete O-GlcNAc proteome. In addition, this work will aid development of O-GlcNAcase inhibitors that target the peptide binding site.

INTRODUCTION

Posttranslational modification of serines/threonines on intracellular eukaryotic proteins with O-linked N-acetylglucosamine (O-GlcNAc) is involved in numerous cellular processes such as transcription, cell cycle progression, and signal transduction (Hart et al., 2007; Love and Hanover, 2005). More than 1,000 proteins are known to be O-GlcNAcylated, and crosstalk with protein phosphorylation is believed to be extensive (Wang et al., 2010). Reversible protein O-GlcNAcylation is achieved by the action of two enzymes, O-GlcNAc transferase (OGT) and O-GlcNAcase (OGA). No precise O-GlcNAcylation sequence motif (sequon) has so far been defined, although promising site mapping data/tools have recently been reported (Chalkley et al., 2009; Wang et al., 2010; Zachara and Hart, 2004). OGT possesses an unusual N terminus, consisting of up to 13.5 tetratricopeptide repeats (TPRs) that are thought to play a role in recognition of intact protein substrates (Clarke et al., 2008; Iyer and Hart, 2003; Lubas and Hanover, 2000). Although recent studies have reported the structure of a bacterial OGT homolog (Clarke et al., 2008; Martinez-Fleites et al., 2008) and the structure of hOGT in complex with a peptide (Lazarus et al., 2011), the molecular mechanisms through which these TPRs contribute to selectivity of O-GlcNAc transfer are not yet understood.

The enzyme that removes O-GlcNAc, OGA, is a 103 kDa enzyme with two domains: an N-terminal hydrolase catalytic domain belonging to CAZy family GH84 (Henrissat and Davies, 1997) and a C-terminal domain that has been proposed to possess histone acetyltransferase activity (Gao et al., 2001; Toleman et al., 2004). Although human OGA (hOGA) can be expressed and purified yielding samples suitable for biochemical studies, attempts to crystallize the protein have so far failed. We recently used an apparent OGA homolog from C. perfringens (CpOGA) to provide insights into the OGA structure (Rao et al., 2006), in parallel with a study on a similar enzyme from B. thetaiotaomicron (Dennis et al., 2006). These structures have identified the active site, which is almost fully conserved with hOGA, and have revealed the molecular details of the interaction with the GlcNAc sugar (Dennis et al., 2006; Rao et al., 2006). Complexes of these enzymes with widely used inhibitors of hOGA have facilitated structure-based design of the potent and selective thiazoline/GlcNAcstatin-based hOGA inhibitors (Dorfmueller et al., 2010; Yuzwa et al., 2008). Strikingly, both these bacterial enzymes were shown to possess O-GlcNAcase activity toward a broad spectrum of O-GlcNAc proteins in human cell lysates (Dennis et al., 2006; Yuzwa et al., 2008). Similar to OGT, it is not clear whether hOGA possesses a (glyco)peptide sequence preference, or how the enzyme binds glycopeptides and catalyzes their O-GlcNAc removal. Thus, the currently available structural information for OGT and OGA does not explain the molecular mechanisms of their interactions with protein substrates, and this limits our understanding of regulation of O-GlcNAc turnover and cycling rates. We investigated how OGA interacts with glycopeptide substrates, revealing that intramolecular interactions in the substrate may affect binding to the OGA active site, and elucidating how OGA achieves O-GlcNAc removal from O-GlcNAc sites in a sequence-independent manner.

RESULTS AND DISCUSSION

O-GlcNAcase Glycopeptide Hydrolysis Is Independent of Peptide Length

Although the structure of human OGA (hOGA) is not available, high-resolution structures of a homologous protein, NagJ, from C. perfringens have been reported, including complexes with the hOGA inhibitors PUGNAc (Rao et al., 2006) and GlcNAcstatin (Dorfmueller et al., 2006). Although it is unclear whether CpOGA
Figure 1. OGA Binds Different O-GlcNAc Glycopeptides with Similar Conformation but Different Affinities

(A) Origin and sequence of O-GlcNAc peptides. The sequence of the longest peptides used is given, with residues observed in the crystal structure highlighted by colored boxes, and the shorter peptides indicated by brackets.
is a physiological OGA, it shows significant in vitro O-GlcNAcase activity on O-GlcNAc proteins in lysates from human cell lines and possesses a putative substrate-binding groove conserved with hOGA (Rao et al., 2006; Schimpl et al., 2010). Here we report that CpOGA, like hOGA (Schimpl et al., 2010), possesses activity toward synthetic glycopeptides derived from validated O-GlcNAc sites in the human proteome, namely p53 (Ser149 [Yang et al., 2006]), TAB1-binding protein 1 (TAB1, Ser295 [Schimpl et al., 2010]), and hOGA itself (Ser405 [Lazarus et al., 2006]) (Figure 1). The Michaelis constants ([OGA] = 10 μM for CpOGA and correlate with the values observed for hOGA (r = 0.91; see Figure S1 available online), suggesting that the bacterial enzyme is a suitable model for understanding hOGA-substrate interactions. Interestingly, there is no correlation between Km for either enzyme and glycopeptide length, suggesting that the range in Km is observed for the different peptides (3–47 μM for CpOGA, 21–630 μM for hOGA; Figure 1B) must stem from structural/sequence properties near the O-GlcNAc site.

OGA Binds Different O-GlcNAc Peptides with Similar Conformations

To investigate the molecular basis of this substrate specificity, we exploited the catalytic acid mutant of CpOGA, D298N, which is inactive yet unaffected in its ability to bind substrate (Rao et al., 2006; Schimpl et al., 2010). We generated an alternative CpOGA crystal form with a highly accessible active site (see Figure S2) and determined the structures of CpOGA D298N in complex with the p53-, TAB1-, and hOGA-derived O-GlcNAc peptides. Synchrotron diffraction experiments resulted in clear unbiased electron density difference maps that defined the conformations of all three glycopeptides (Figure 1C). These complexes define the molecular basis of how OGA recognizes both the sugar and protein components of physiologically relevant substrates. The O-GlcNAc sugar occupies the same position in all three structures (maximum atomic shift of 0.2 Å). It is tethered by extensive hydrogen bonding with residues that are identical between CpOGA and hOGA (Figure 1D) and adopts the 1,2-D face conformation predicted from mechanistic studies and observed in complexes with pseudosubstrates (Macauy and Voadal, 2010; Macauy et al., 2005; Rao et al., 2006). Compatible with the proposed substrate-assisted catalytic mechanism, the carbonyl oxygen approaches the anomeric carbon to within 3.0 Å, poised for nucleophilic attack and in-line displacement of the glycosidic oxygen (angle of 164°).

Glycopeptide Substrate Recognition Involves Backbone Contacts

Strikingly, the backbone of all three glycopeptides run in the same direction (Figure 1D), together defining the −4 through +3 subsites, and adopt similar conformations near the O-GlcNAc site (maximum Cα shift of 1.8 Å for the −2 to +1 subsites). Notably, all side chains point away from the surface of the enzyme (apart from Trp146 in the p53 peptide, which appears to stack with Asn298 of CpOGA D298N), explaining how a single OGA enzyme is able to recognize >1,000 O-GlcNAc proteins. Hydrophobic stacking of the solvent-exposed Tyr189 (Tyr69 in hOGA) aromatic side chain with the −1 and −2 peptide bonds contributes a major amount (~30% of the buried surface) of the interaction between the enzyme and the peptide component of the glycopeptides (Figure 1D). To test the contribution of this interaction to substrate binding, we mutated the corresponding Tyr69 in the human enzyme to Ser, Lys, Gln, and Phe and determined the Km values for several substrate O-GlcNAc peptides (Table 1). Only the Y69F mutant shows no loss in activity, indicating that the hydrophobic stacking interaction is essential for OGA activity.

Table 1. Hydrophobic Stacking Interactions of Tyr69 Are Essential for Substrate Binding by hOGA

| Glycopeptide | TAB1 Peptide | hOGA Peptide | p53 Peptide | p53 Peptide |
|--------------|--------------|--------------|-------------|-------------|
| Y69S         | >4,000       | >4,000       | 3,400 ± 500 | 290 ± 20    |
| Y69K         | >4,000       | >4,000       | >4,000      | 350 ± 20    |
| Y69Q         | 3,600 ± 300  | >4,000       | >4,000      | 170 ± 40    |
| Y69F         | 1,100 ± 60   | 1,200 ± 110  | 2,700 ± 400 | 53 ± 5      |
| Wild-type    | 940 ± 80     | 790 ± 40     | 1,300       | 21 ± 3      |

Tyr69 in human OGA may participate in hydrogen bonding with the catalytic aspartate (Asp175) as well as providing a stacking platform for the −1/−2 peptide bond in the substrate. The Y69S mutation disrupts both these interactions, the Y69K and Y69Q mutants cannot participate in pi-pi stacking, and Y69F abolishes the hydrogen bond. Point mutations were introduced in hOGA, and the Michaelis constant (Km, given in μM) was determined for several peptide substrates.

*Schimpl et al., 2010.

(B) Km of glycopeptide substrates for hOGA (displayed by filled bars [values for the three shortest hOGA- and TAB1-derived peptides reproduced from Schimpl et al., 2010]) and CpOGA (unfilled bars, right y axis) as determined by substrate competition assay with error bars representing the error of curve-fit (see Experimental Procedures for details).

(C) Conformation of O-GlcNAc peptides bound in the active site of CpOGA as determined by X-ray crystallography. Peptides are shown as sticks with colored carbons (TAB1 peptide, yellow; hOGA peptide, green; p53 peptide, orange), with the GlcNAc sugar highlighted by pink carbons. Unbiased [Fo − Fc]/σ(Fo) electron density (i.e., before addition of any glycopeptide model) is shown in gray (contoured at 2.2σ). Intramolecular hydrogen bonds are shown by dashed lines.

(D) O-GlcNAc peptides (sticks) in complex with CpOGA D298N (surface representation). Sequence conservation between hOGA and CpOGA is indicated by blue shading of identical residues on the molecular surface. The catalytic acid Asp298 (mutated to Asn) and the conserved Tyr198 are labeled, with corresponding residue numbers for hOGA given in brackets.

See also Figures S1 and S2.
Table 2. Intramolecular Hydrogen Bonds Stabilize the Substrate Conformation and Reduce the $K_m$ for hOGA

| Original sequence | TAP1 Peptide (Ser395) | p53 Peptide (Ser149) |
|------------------|-----------------------|----------------------|
| $K_m$ (hOGA)     | 4100 ± 300 µM         | 21 ± 2 µM            |
| Altered sequences| VPYS(O-GlcNAc)SAQ    | QLVD(S-O-GlcNAc)TPPPG|
| $K_m$ (hOGA)     | 810 ± 140 µM          | 3700 ± 900 µM        |
|                  | QLVS(O-GlcNAc)TPPPG   |                      |
| $K_m$ (hOGA)     | 1800 ± 200 µM         |                      |

The contribution of intramolecular hydrogen bonds toward substrate binding was probed by introducing a potential hydrogen bond donor in the sequence of the TAP1 peptide. For the p53 peptide, the hydrogen bond was disrupted by replacing either the donor or the acceptor with isosteric aliphatic amino acids (highlighted in boldface and underscored).

Intramolecular Hydrogen Bonds Affect Substrate Conformation and $K_m$

All three glycopeptides adopt a “V-shaped” conformation that allows the sugar to penetrate the OGA active site. Interestingly, for two of the peptides, this conformation appears to be stabilized by intramolecular hydrogen bonds (Figure 1C). The hOGA-derived glycopeptide forms a hydrogen bond between the histidine in the –1 subsite and the backbone carbonyl oxygen of the O-GlcNAc serine (Figure 1C). For the p53-derived glycopeptide, a hydrogen bond is observed between the aspartic acid in the –1 subsite and the threonine in the +1 subsite (Figure 1C). Such intramolecular interactions may stabilize the OGA-bound conformation of these glycopeptides, explaining the significantly lower $K_m$'s compared to the TAP1-derived peptide. We tested this hypothesis by designing glycopeptide sequences that either disrupted (in case of the p53-derived peptide) or introduced (in case of the TAP1-derived peptide) such intramolecular interactions (see Table 2). Indeed, introducing a hydrogen bond acceptor in the TAP1 peptide leads to a 5-fold decrease in $K_m$ for hOGA, whereas disrupting the hydrogen bond in the p53-derived peptide, either by removing the hydrogen bond donor or acceptor, leads to an 85-fold increase in $K_m$. Thus, although OGA substrate recognition does not appear to involve direct recognition of specific residues proximal to the O-GlcNAc site, subtle conformational effects appear to tune substrate recognition. It is possible that sequence-dependent stabilization of a specific backbone conformation around the O-GlcNAc site could give rise to differential persistence/cycling rates for individual O-GlcNAc sites in the human proteome.

O-GlcNAcylation of p53 Involves Limited Conformational Change around the Acceptor Serine

There are currently no available structures of O-GlcNAc glycoproteins, and O-GlcNAc-sites on structurally characterized proteins appear to reside in disordered/structurally undefined regions, limiting our understanding of the conformational changes induced by protein O-GlcNAcylation, or how these proteins would interact with OGA/OGT. A notable exception is the tumor suppressor protein p53, where the reported Ser149 O-GlcNAc site resides in a loop that is fully defined in the crystal structure of the p53 DNA binding domain (Cho et al., 1994). Ser149 lies at the tip of this loop, projecting out into the solvent. Comparing the structure of this loop in the p53 DNA binding domain structure with the glycosylated form reported here, it is apparent that the overall trajectory of the loop is approximately conserved between the two conformations of the peptide (Figure 2A; average Cα shift of 2.2 Å, although a number of side chain flips are observed (in particular Trp146). This superposition can also be expanded to the CpOGA and p53 proteins, yielding a model of an O-GlcNAcase-glycoprotein substrate complex (Figure 2B), with p53 and its Ser149 loop occupying the OGA putative substrate binding site. Further work will be needed to establish how regions beyond the immediate vicinity of the O-GlcNAc site will contribute to the interaction of OGA with the full p53 DNA binding domain.

Different OGA Inhibitors Display Varying Levels of Peptide Mimicry

The three most potent and widely used OGA inhibitors are the transition state mimics PUGNac (Haltiwanger et al., 1992) and GlcNAcstatin (Dorfmueller et al., 2010), as well as the NAG-thiazoline derivatives that mimic the oxazoline reaction intermediate (Macauley et al., 2005; Yuzwa et al., 2008). Thiazolines, such as Thiamet-G, only occupy the sugar pocket, whereas PUGNac and GlcNAcstatin contain additional substituents mimicking the aglycon. Comparison with the CpOGA-glycopeptide complexes reported here reveals that these phenyl moieties in fact occupy the +1/+2 subsites accommodating the glycopeptide backbone (Figure 2C). Since the use of PUGNac and the thiazolines in probing the role of O-GlcNAc in modulating insulin sensitivity has yielded contradictory results (Macauley et al., 2008; Vosseller et al., 2002), further investigation is required to establish whether the different binding modes of these O-GlcNAcase inhibitors may explain the discrepancies between their effects in vitro.

Conclusions

To our knowledge, this work reports the first structures of an enzyme of the O-GlcNAcase family in complex with glycopeptide substrates. Despite carrying different sequences, the glycopeptides adopt similar conformations in the active site. Crucially, while O-GlcNAcase does interact with the glycopeptide substrate backbone through specific enzyme side chains, the glycopeptide substrate side chains face away from the enzyme’s binding cleft, explaining how a single enzyme can target a plethora of O-GlcNAc proteins. However, specific intramolecular interactions in the glycopeptide may predispose certain amino acid sequences for a conformation that is compatible with the O-GlcNAcase binding cleft. This work will underpin a mechanistic interpretation of differential cycling rates of sites in the O-GlcNAc proteome and facilitate development of inhibitors that not only target the sugar binding pocket, but also the peptide binding groove.

SIGNIFICANCE

Protein O-GlcNAcylation is an essential and reversible glycosylation event in higher eukaryotes, where hundreds of intracellular proteins are O-GlcNAcylated. O-GlcNAc
addition and removal is catalyzed by O-GlcNAc transferase and O-GlcNAcase, respectively. How a single pair of enzymes achieves cycling of the complete O-GlcNAc proteome is one of the key questions in the field. We report the molecular details of the interaction of a bacterial O-GlcNAcase and three synthetic O-GlcNAc peptides matching established O-GlcNAc sites in the human proteome. In the 3D structures, we observe recognition of the sugar moiety as well as sequence-independent interactions with the peptide backbone, thus elucidating the molecular basis of the broad substrate specificity of the O-GlcNAcase enzyme. We report some influence of the peptide sequence directly surrounding the modification site; intramolecular hydrogen bonding within the peptide facilitates the binding to the enzyme. Peptides capable of forming such interactions are better O-GlcNAcase substrates in vitro, and we hypothesise that the cycling rate of individual O-GlcNAc sites in vivo may vary depending on the surrounding protein sequence. Finally, this work will aid development of O-GlcNAcase inhibitors that target the peptide binding site.

**Experimental Procedures**

**Glycopeptide Synthesis**

Microwave-assisted solid phase peptide synthesis was performed with a CEM Liberty automated peptide synthesizer on low load Rink amide MBHA resin 100-200 mesh (Novabiochem) using standard Fmoc chemistry protocols on a 0.05 mmol scale. The 3,4,6-triacetyl-O-GlcNAc-Fmoc-Ser-OH building block was synthesized as described previously (Schimpl et al., 2010). All peptides were N-terminally acetylated and C-terminally amidated, and were purified via high-performance liquid chromatography.

**hOGA Expression and Purification**

The coding sequence for full-length human OGA was cloned into pEBG6P. Mutations were introduced using the following oligonucleotide primers:

\[
\begin{align*}
618 \text{stop} & \quad 5'\text{-caagaagcttaaaggtggatcaattgaattagtgtaaattg} & 5'\text{-catatttgcgaacttc}
\end{align*}
\]

hOGA glycopeptide hydrolysis assays were carried out as described previously (Dorfmueller et al., 2006; Schimpl et al., 2010) using multisubstrate enzyme kinetics with the fluorogenic pseudosubstrate 4MU-GlcNAc as the reporter substrate. Briefly, initial rates of hydrolysis of 4MU-GlcNAc were determined in the presence of increasing concentrations of glycopeptide, and the Michaelis constant of the competing substrate (\(K_m\)) was determined using the following equation:

\[
\frac{v}{v_t} = \frac{1 + K_m}{S} \left(1 + \frac{K_m}{S} \right)
\]
where $v/v_0$ is the relative activity in the presence of inhibitor, $K_i$ and $S$ are the Michaelis constant and substrate concentration of the reporter substrate, and $S'$ is the concentration of glycopeptide. Reactions were performed at 37°C in 50 mM citrate-phosphate buffer (pH 7.4) and 0.1 mg/ml BSA. Experiments were performed in triplicate, and data were analyzed and plotted with GraphPad PRISM.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes two figures and one table and can be found with this article online at doi:10.1016/j.chembiol.2012.01.011.

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