Molecular and Structural Characterization of Hexameric β-D-Glucosidases in Wheat and Rye

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The wheat (Triticum aestivum) and rye (Secale cereale) β-D-glucosidases hydrolyze hydroxamic acid-glucose conjugates, exist as different types of isozyme, and function as oligomers. In this study, three cDNAs encoding β-D-glucosidases (TaGlu1a, TaGlu1b, and TaGlu1c) were isolated from young wheat shoots. Although the TaGlu1s share very high sequence homology, the mRNA level of Taglu1c was much lower than the other two genes in 48- and 96-h-old wheat shoots. The expression ratio of each gene was different between two wheat cultivars. Recombinant TaGlu1b expressed in Escherichia coli was electrophoretically distinct from TaGlu1a and TaGlu1c. Furthermore, coexpression of TaGlu1a and TaGlu1b gave seven bands on a native-PAGE gel, indicating the formation of both homo- and heterohexamers. One distinctive property of the wheat and rye glucosidases is that they function as hexamers but lose activity when dissociated into smaller oligomers or monomers. The crystal structure of hexameric TaGlu1b was determined at a resolution of 1.8 Å. The N-terminal region was located at the dimer-dimer interface and plays a crucial role in hexamer formation. Mutational analyses revealed that the aromatic side chain at position 378, which is located at the entrance to the catalytic center, plays an important role in substrate binding. Additionally, serine-464 and leucine-465 of TaGlu1a were shown to be critical in the relative specificity for DIMBOA-glucose (2-O-β-D-glucopyranosyl-4-hydroxy-7-methoxy-1,4-benzoxazin-3-one) over DIBOA-glucose (7-demethoxy-DIMBOA-glucose).

β-D-Glucosidase (EC. 3.2.1.21) is a major member of the family GH1 and GH3 glycoside hydrolases and is responsible for hydrolysis of terminal non-reducing β-D-Glc residues in oligosaccharides (or polysaccharides) or glucoconjugates. In plants, β-D-glucosidases are involved in various functions, including lignification (Dharmawardhana et al., 1995), regulation of the biological activity of cytokinins (Brzobohatý et al., 1993; Falk and Rask, 1995; Haberer and Kieber, 2002), control of the biosynthesis of indole-3-acetic acid (Ljung et al., 2001; Persans et al., 2001), and chemical defense against pathogens and herbivores (Niemeyer, 1988; Sicker et al., 2000; Zagrebelyn et al., 2004). Many secondary products in plants occur as glucosconjugates with one or two Gc units attached to a hydroxy or thiol group. Hydrolysis of the glucosidic linkage in secondary metabolites, such as cyanogenic-, flavonoid-, and hydroxamic acid-glucosides, can drastically alter the biological activity, chemical stability, and water solubility of the molecule. The β-D-glucosidases implicated in the hydrolysis of plant secondary metabolites are members of the family GH1 glycoside hydrolases. The classification system of the glycoside hydrolases is available on the CAZy database at http://afmb.cnrs-mrs.fr/CAZY/.

Although β-D-glucosidases possess broad substrate specificity with respect to the aglycone moiety, the preferred aglycone structures vary with each glucosidase, reflecting their wide variety of physiological roles. Indeed, some β-D-glucosidases exhibit strict aglycone specificity. For example, the sorghum (Sorghum bicolor) glucosidase (dhurrinase 1, SbDhr1) acts specifically on its natural substrate, dhurrin. Dhurrin inhibits the activity of the maize (Zea mays) homolog, ZmGlu1, whose amino acid sequence shares about 70% identity with SbDhr1 (Hösel et al., 1987; Cicek and Esen, 1998). Recently, several research groups have investigated aglycone recognition mechanisms by SbDhr1 and ZmGlu1 using a combination of site-directed mutagenesis and x-ray crystallography (Czizek et al., 2000, 2001; Verdoucq et al., 2003, 2004).

In previous studies, we purified β-D-glucosidases from the seedlings of wheat (Triticum aestivum) and rye (Secale cereale; Sue et al., 2000a, 2000b). The seedlings accumulate O-β-D-glycosides of hydroxamic acids (HxS; 2,4-dihydroxy-1,4-benzoxazin-3-one, DIBOA, and its 7-methoxy derivative, DIMBOA; Fig. 1) as defensive compounds against pathogens and herbivores.

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[The online version of this article contains Web-only data.]
multiple activity bands (Sue et al., 2000b). The N-terminal different ratios, which is observed in a zymogram as several types of heterooligomers by assembling in subunits in the wheat enzyme are thought to make up homodimer of 60-kD subunits. Furthermore, the two

This is different from the maize homolog that is a hexameric; Sue et al., 2000, 2001; Zouhar et al., 2001) and are thought to exist in an oligomeric form (probably tetrameric or octameric quaternary structure, and the influence of oligomerization on activity are not yet known. The crystal structures of four family GH1 glucosidases from plants have been solved (Barrett et al., 1995; Burmeister et al., 1997; Czjzek et al., 2001; Verdoucq et al., 2004), and biochemical analysis has shown these enzymes function as dimers. Some bacterial enzymes have shown tetrameric or octameric quaternary structure in the crystal structure (Aguilar et al., 1997; Sanz-Aparicio et al., 1998; Chi et al., 1999; Hakulinen et al., 2000). However, the amino acid sequence of rye glucosidase, which forms an oligomer, shows a higher level of similarity to the plant enzymes than to the bacterial enzymes. Additionally, the monomer-monomer interaction and orientation of the bacterial glucosidases are different from those of the plant enzymes ZmGlu1 and SbDhr1. Thus, it is of interest to elucidate the quaternary structure of wheat glucosidase and investigate the structure-activity relationships of this enzyme. In this study, we cloned three cDNAs encoding the wheat 

Primary Structure of the Wheat \( \beta \)-D-Glucosidase

Three cDNAs encoding wheat \( \beta \)-D-glucosidases were isolated by screening a cDNA library prepared from 48-h-old wheat shoots (cv Chinese Spring [CS]) and designated \( \text{Taglu1a}, \text{Taglu1b}, \) and \( \text{Taglu1c} \) (supplemental text named as “Cloning of the wheat glucosidases”). \( \text{Taglu1a}, \text{Taglu1b}, \) and \( \text{Taglu1c} \) comprised open reading frames of 1,710-, 1,710-, and 1,713-bp encoding polypeptides of 569, 569, and 570 amino acids, respectively (Supplemental Fig. 1). The deduced amino acid sequence of \( \text{Taglu1a} \) shows 91%, 95%, and 95% identity and 95%, 98%, and 97% similarity to ScGlu, TaGlu1b, and TaGlu1c, respectively. Each TaGlu1 included the N-terminal sequence(9,10),(989,992)

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Transcript Profiles of the Genes Encoding TaGlul
Isozymes in Young Wheat

In our previous work, we reported that the glucosidase activity changes transiently, peaking 36 to 48 h after imbibition (Sue et al., 2000b). Northern-blot analysis showed that the genes are expressed at a high level 36 to 48 h after imbibition and that expression level then gradually decreases as the plant grows (Fig. 2A). This pattern correlated well with that of glucosidase activity. However, northern-blot analysis could not discriminate between each of the three Taglu1 genes because of the high level of sequence homology. Therefore, the expression of each glucosidase gene was analyzed by quantitative PCR using primers specific to each gene. In this experiment, RNA was also prepared from another bread wheat cultivar, Asakazekomugi (Ak), in addition to CS, to examine whether the expression pattern of each gene is conserved among the two cultivars.

In CS, Taglu1a was most highly expressed (67%) in 48-h-old wheat, with Taglu1b expressed at about one-half this level (32%; Fig. 2B). The ratio was slightly different in a 96-h-old plant where Taglu1a expression increased to 85% and Taglu1b decreased to 14%, though the total amount of glucosidase gene expression declined as shown by the northern analysis. In both growth stages, the expression levels of Taglu1c were much less (0.4% and 0.1% in 48- and 96-h-old plants, respectively) than those of other glucosidase genes. These data were supported by the results of the cDNA library screening; 37 clones of Taglu1a, 11 clones of Taglu1b, and only one clone of Taglu1c were obtained. In contrast to the results of CS, Taglu1a and Taglu1b (Fig. 2B) were shown to be expressed in almost equal amounts in 48-h-old Ak (Taglu1a, 51%; Taglu1b, 48%). As the plant grows, however, the expression of Taglu1a and Taglu1b changed to a percentage comparable to that in CS. Similarly to CS, Taglu1c was expressed at a low level in both 48- and 96-h-old AK plants (1.3% and 4.7%, respectively).

Figure 2. Expression level of Taglu1 mRNA. A, Northern-blot analysis. Taglu1, probed with Taglu1a; rRNA, ribosomal RNA stained with ethidium bromide. B, Quantification of Taglu1a to Taglu1c mRNA in 48- and 96-h-old wheat shoots (CS and Ak). The data of each growth stage are described as the ratio of each gene to the sum of Taglu1a to Taglu1c.
peaks corresponding to the dimer and tetramer were occasionally observed (data not shown). Thus, the wheat glucosidase also can exist as a dimer and a tetramer. However, the fractions containing these oligomers, as well as the monomer, showed only slight activity with DIMBOA-Glc. The activity was so low that we could not eliminate the possibility that the activity was derived from minor contamination with the active hexamer. When the purified hexameric glucosidase was dialyzed against 50 mM HEPES without NaCl, the monomer readily formed (Fig. 4B), resulting in loss of activity.

Among the isozymes, TaGlu1b showed the highest activity with DIBOA-Glc and DIMBOA-Glc (the $k_{cat}/K_m$ values 149 and 4,138 s$^{-1}$/mM, respectively), while TaGlu1a, the major isozyme in 48-h-old CS, showed the lowest activity (4.5-fold lower than TaGlu1b). The lower activity of TaGlu1a may be caused by instability of the active hexamer, as suggested by gel filtration where the amount of hexameric TaGlu1a was lower than that of TaGlu1b or TaGlu1c (data not shown).

Figure 3. SDS- and native-PAGE of the wheat and rye glucosidases. A, The glucosidase monomers were analyzed by SDS-PAGE. For the native (without His-tag) glucosidases, the crude E. coli cell extracts were directly subjected to SDS-PAGE. The His-tagged glucosidases were electrophoresed after purification by affinity chromatography on a nickel-charged column. B, The cell extracts were subjected to native-PAGE (on an 8% separating gel for 4 h) and stained with Coomassie Brilliant Blue. The arrowheads indicate the seven types of glucosidase hexamers expressed in coexpression lines. C, The bands with $\beta$-glucosidase activity were detected by activity staining. The crude extracts of E. coli cells and 48-h-old shoots were separated under nondenaturing conditions. In each segment, M, a, b, c, a/b, b/c, W, and R indicate marker proteins, TaGlu1a, TaGlu1b, TaGlu1c, coexpressed TaGlu1a and TaGlu1b, coexpressed TaGlu1b and TaGlu1c, wheat shoots, and rye shoots, respectively.

Figure 4. Gel filtration of TaGlu1a, ScGlu, and Zm-Glu1a. A, The enzyme solution eluted from the nickel column was further purified by gel filtration using a Superdex 200 column. Each fraction volume was 0.5 mL. The eluted protein was monitored by $A_{280}$. The two protein peaks correspond to hexamer and monomer. B, Solid line, Gel filtration of ScGlu was performed after affinity chromatography. Two peaks of hexamer and monomer were observed. Dashed line, TaGlu1a (wild type) purified by affinity chromatography followed by gel filtration was dialyzed against HEPES buffer without NaCl and subjected to gel filtration analysis. Only a monomeric protein was detected. Dash-dot line, Zm-TaGlu1a purified by affinity chromatography was further purified by gel filtration. While the hexamer peak was not detected, the dimer and monomer peaks were observed.
Hexameric \( \beta\)-D-Glucosidases from Wheat and Rye

Crystal Structure of the Wheat \( \beta\)-D-Glucosidase and the Role of Its N-Terminal Region in Hexamer Formation

The structure of TaGlu1b in complex with DIMBOA was determined at 1.8-Å resolution from a crystal that was soaked in the DIMBOA solution. The final refinement statistics are shown in Table I. The overall structure of TaGlu1b was almost the same as that of known \( \beta\)-D-glycosidases, which have classical (\( \beta/\alpha \))\(_6\) barrel folds. The \( \beta\)-strands and \( \alpha\)-helices within each \( \beta/\alpha \) repeat were connected by loops at the top barrels. One disulfide bond was observed between Cys-210 and Cys-216, which is conserved among \( \beta\)-D-glycosidases in sorghum and maize. Although 11 residues at the N terminus and 18 at the C terminus of the mature enzyme were not included in the structure due to the lack of electron density, both termini were modeled three residues longer than those of sorghum or maize structures. The program PROCHECK (Laskowski et al., 1993) was applied to validate the structure; 374 residues were in most favored, 48 in additional allowed, and two (Ala-78 and Trp-463) were in generously allowed regions. When we superposed the TaGlu1b structure with those from sorghum (PDB code 1V03) and maize (PDB code 1E4N), they mostly fitted well, except that slightly different conformations were observed in the region from Val-413 to Pro-422 (loop D; Fig. 5C), probably due to low sequence similarity (Supplemental Fig. 1). The electron density of the amino acid chain was clearly observed, whereas that of the aglycone was not defined. This may have been due to interference caused by several glycerol molecules bound in the active site. The binding of a glycerol molecule at the active site was reported for the maize glucosidase Zm-p60.1 (Zouhar et al., 2001). A sulfate ion derived from LiSO\(_4\) in the crystallization buffer was also observed, which was fixed by Ser-366 and Asp-271, and Arg-434 of an adjacent subunit as well. Although the asymmetric unit contained one monomer of the enzyme, the symmetrical operation produced a hexamer conformation (Fig. 5A), where the dimer is obtained by the crystallographic 2-fold symmetry operation of the monomer and the hexamer by the 3-fold symmetry operation of the dimer. Both symmetrical operation axes are located perpendicular to each other.

If we regard the hexamer as three molecules of dimer (equivalent to the maize and sorghum dimers), the N-terminal region of TaGlu1b is located at the interface between the adjacent dimers. With respect to this region, the subunits are linked by four direct hydrogen bonds (Fig. 5B). To examine the role of this region in the formation of hexameric structure, the N-terminal 25 residues of the mature TaGlu1a and TaGlu1b were replaced with the corresponding residues of ZmGlu1 (Supplemental Fig. 1), considering that this enzyme is known to exist as a dimer. The chimeric glucosidases (Zm-TaGlu1a and Zm-TaGlu1b) completely lost their ability to form a hexamer, which was confirmed by gel filtration chromatography (Fig. 4B). Instead of being hexameric, the dimeric structure was the major component on the chromatogram, suggesting the crucial role of the N-terminal sequence in maintaining the dimer-dimer association. The fractions containing dimeric Zm-TaGlu1a or Zm-TaGlu1b exhibited little activity toward DIMBOA-Glc (data not shown).

Site-Directed Mutagenesis of the Substrate Binding Pocket

The recombinant enzyme with an N-terminal Histag displayed an activity comparable to the naturally occurring glucosidase; the \( V_{\text{max}} \) value of the natural wheat glucosidase for DIMBOA-Glc was 4,100 nkat/mg protein (Sue et al., 2000b) and that of the recombinant TaGlu1a was 5,200 nkat/mg protein. We therefore used the N-His-tagged enzyme for site-directed mutagenesis of residues at the substrate binding pocket. The amino acid residues involved in the substrate binding pocket are absolutely conserved among TaGlu1a to TaGlu1c (Supplemental Fig. 1), deduced from the structural and the sequence alignments with ZmGlu1-E191D mutant in complex with DIMBOA-Glc (Czjzek et al., 2000). The data of enzyme activity are shown in Tables II and III.

The primary structures of the aglycone binding sites of TaGlu1a and ScGlu diverge from each other at the residues Ser-464 and Leu-465 in TaGlu1a and Gly-464 and Ser-465 in ScGlu. While the mutations G464S and S465L of ScGlu decreased the relative efficiency for DIBOA-Glc by 42% and 62%, respectively, they increased the efficiency for DIMBOA-Glc by 100% and 143%, respectively. The effects were enhanced by introduction of the double mutation G464S/S465L. Introduction of Phe at position 464 of both enzymes resulted in decreased efficiency for the natural substrates. The influence was most obvious in TaGlu1a with

### Table I. Refinement statistics

\[
\begin{align*}
R & = \frac{\Sigma |F_{\text{obs}}| - \Sigma |F_{\text{calc}}|}{\Sigma |F_{\text{obs}}|} \\
R_{\text{free}} & = \frac{\Sigma |F_{\text{obs}}| - \Sigma |F_{\text{calc}}|}{\Sigma |F_{\text{obs}}|} \\
R_{\text{free}} & = \frac{5\%}{100\%}
\end{align*}
\]

<table>
<thead>
<tr>
<th>TaGlu1b</th>
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<td>Space group</td>
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<tr>
<td>Cell dimensions</td>
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</tr>
<tr>
<td>Resolution (Å)</td>
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<td>No. reflections</td>
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<td>Completeness (%)</td>
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<tr>
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<td>Luzzati ESD (obs)</td>
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<tr>
<td>Luzzati ESD (R(_{\text{free}}))</td>
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</tr>
<tr>
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<td>Bond lengths (Å)</td>
<td>1.3</td>
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DIMBOA-Glc (relative efficiency 5.5%). The $k_{cat}/K_m$ values of ScGlu-F198A for DIMBOA-Glc and DIBOA-Glc decreased by a factor of 24- to 28-fold as compared with those of wild-type ScGlu. However, the decrease in the $k_{cat}/K_m$ value of this mutant was solely due to an increased $K_m$, whereas the $k_{cat}$ value actually increased. TaGlu1a-F198A mutants showed some activity ($k_{cat}$) toward the natural substrates, although the $K_m$ values increased dramatically. TaGlu1a-F198A mutants showed some activity ($k_{cat}$) toward the natural substrates, although the $K_m$ values increased dramatically. The Y378A mutation of ScGlu enhanced $k_{cat}/K_m$ values for DIMBOA-Glc and pNP-Glc by 300% to 380%, whereas that of TaGlu1a was lowered to 11% to 35% for all substrates tested. Although the catalytic efficiency ($k_{cat}/K_m$) of the ScGlu mutant for DIBOA-Glc was comparable to that of wild type, both the $K_m$ and $k_{cat}$ were about 10-fold greater. Replacing Phe-471 in TaGlu1a with Tyr decreased the catalytic efficiency for all the substrates tested. However, the same mutation on ScGlu increased the $k_{cat}$ for the three substrates by about 100% and decreased $K_m$ for DIMBOA-Glc and pNP-Glc. TaGlu1 and ScGlu contain the well-conserved TFNEP and ITNEG motifs at the catalytic center of family GH1 glucosidases, and replacement of either of the two Glu residues in the motifs (Glu-191 and Glu-407, respectively) resulted in a complete loss of enzyme activity.

DISCUSSION

TaGlu1a and TaGlu1b Genes Are Highly Expressed in Young Wheat Shoots

We isolated three genes encoding family GH1 glucosidases that are responsible for the hydrolysis of...
The transcript profile of Taglu1 genes, as analyzed by northern hybridization, agreed with the transient occurrence of glucosidase activity and Hx-Glc occurrence in young plants (Sue et al., 2000b). However, the expression profiles of each gene after 48- and 96-h imbibition do not appear to be synchronized, suggesting that the expression of these three genes is controlled independently of each other. This is similar to the expression profiles of the three homoeologous genes of each of the five Hx biosynthetic genes, which vary between developmental stages of wheat seedlings (Nomura et al., 2005). The mechanism underlying the different expression profiles of the three Taglu1s as well as the biosynthetic genes has not yet been uncovered.

The ratio of each Taglu1 gene differs between two wheat cultivars at the same growth stage. At 48 h, CS expressed the Taglu1a gene at a higher ratio than Taglu1b, whereas Ak expressed both genes almost equally. In consideration of the result that TaGlu1b moves more slowly than TaGlu1a and TaGlu1c on a native-PAGE gel, the higher expression level of TaGlu1b is presumably due to its more substantial net negative charge compared with TaGlu1a and TaGlu1c.

Table II. Kinetic parameters of the TaGlu1s and TaGlu1a mutants

The relative efficiency toward each substrate is the percent ratio of the $k_{cat}/K_m$ values to that of ScGlu. n.d., Not detected. –, Not determined.

<table>
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<tr>
<th>Substrate</th>
<th>ScGlu</th>
<th>E191A</th>
<th>F198A</th>
<th>Y378A</th>
<th>Y378F</th>
<th>E407A</th>
<th>S464F</th>
<th>Y378A</th>
<th>F471Y</th>
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<td>0.80</td>
<td>n.d.</td>
<td>44.1</td>
<td>7.99</td>
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<td>n.d.</td>
<td>1.60</td>
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<td>0.83</td>
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<td>118</td>
<td>n.d.</td>
<td>233</td>
<td>1,098</td>
<td>12.0</td>
<td>n.d.</td>
<td>54.6</td>
<td>86.9</td>
<td>46.6</td>
</tr>
<tr>
<td>$k_{cat}/K_m$</td>
<td>148</td>
<td>n.d.</td>
<td>5.28</td>
<td>137</td>
<td>23.1</td>
<td>n.d.</td>
<td>34.1</td>
<td>85.2</td>
<td>56.1</td>
</tr>
<tr>
<td>Relative Efficiency</td>
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<td>100</td>
<td>3.6</td>
<td>92.6</td>
<td>15.6</td>
<td>0</td>
<td>23.0</td>
<td>57.6</td>
<td>37.9</td>
</tr>
</tbody>
</table>

Table III. Kinetic parameters of ScGlu and its mutants

The relative efficiency toward each substrate is the percent ratio of the $k_{cat}/K_m$ values to that of ScGlu. n.d., Not detected.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>ScGlu</th>
<th>E191A</th>
<th>F198A</th>
<th>Y378A</th>
<th>Y378F</th>
<th>E407A</th>
<th>S464F</th>
<th>Y378A</th>
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<td>DioBA-Glc</td>
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<td>148</td>
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</table>

The purified glucosidase from 48-h-old Ak showed seven bands on a native-PAGE gel. Because TaGlu1c was considered to be expressed at a much lower level than TaGlu1a and TaGlu1b in the plant, the natural glucosidase would mainly exist as homo- and heterooligomers of TaGlu1a and TaGlu1b.

The Wheat and Rye Glucosidases Function as a Hexamer

The recombinant TaGlu1b showed greater electrophoretic mobility than TaGlu1a and TaGlu1c on a native-PAGE gel. The mobility of proteins in native-PAGE is influenced by the shape, or holding, and charge of the proteins as well as their Mr. The results of ion-spray mass spectrometric analyses of recombinant TaGlu1a and TaGlu1b subunits excluded the possibility of proteolytic digestion during cell disruption. Furthermore, the overall holding of the TaGlu1b must be almost identical with the other isozymes because it shares high homology with TaGlu1a and TaGlu1c with respect to the primary structures and has similar enzyme activity and substrate specificity. Therefore, the greater mobility of TaGlu1b is presumably due to its more substantial net negative charge compared with TaGlu1a and TaGlu1c. This hypothesis is supported...
by the lower theoretical pI value of TaGlu1b (pI 5.2) in comparison with that of TaGlu1a and TaGlu1c (both with pI 5.8). Nevertheless, the anomalously low mobility of TaGlu1b under denaturing conditions cannot be predicted from the pI value. There may be additional factors that cause the mobility discrepancy by SDS-PAGE. The slight differences in the amino acid composition might result in different migration rates for the protein even under denaturing condition.

In the previous study, we demonstrated that the wheat glucosidase exhibits activity as an oligomer (Sue et al., 2000b), but the number of subunits constituting the oligomer remained unclear. These results from coexpression of two TaGlu1 isozymes (TaGlu1a and TaGlu1b) and the corresponding gel filtration analysis have established that TaGlu1 exhibits its activity only as a hexamer. Because wheat and rye glucosidases show similar gel filtration and electrophoretic profiles, it is reasonable to assume that the active form of ScGlu is also a hexamer. Although only three bands were detected in the sample of natural ScGlu on a native-PAGE gel (Fig. 3C), they must comprise a larger number of stacking bands because the rye glucosidase isolated from shoots separated into at least six protein peaks by anion-exchange chromatography (Sue et al., 2000a). Presumably, the rye used in our studies contains multiple isozymes whose electrophoretic mobility is very similar to each other. The maize glucosidase gene (Zmglu1) is known to be a highly polymorphic gene, and a zymogram using SDS-PAGE. The slight differences in the amino acid sequences of the N-terminal region are not make up a hexamer but a dimer (Fig. 4B). The amino acid sequences of the N-terminal region are least conserved among the family GH1 β-D-glucosidases. This result suggests a novel function of the N-terminal region in maintaining the quaternary structure of the β-D-glucosidases. The dimer of Zm-TaGlu1 easily dissociated into monomers (data not shown), suggesting that the monomer of TaGlu1 interacts very weakly in comparison with the ZmGlu1 or SbDhr1 monomers. These characteristics differ from the monocot family GH1 β-D-glucosidases, where the dimeric forms represent the stable structural units from which the oligo- and multimers are composed. The crystal structure of the TaGlu1b monomer resembles the known structures of the maize and sorghum homologs (Fig. 5C). The two residues in ZmGlu1 (Arg-295 and Asp-342) that were shown to form the intermonomer salt bridges (Czjzek et al., 2001) are conserved in TaGlu1b (Arg-296 and Asp-343). The intramolecular disulfide bond responsible for maintaining the dimeric form of Zm-p60.1 (Zouhar et al., 2001) is also conserved in the wheat glucosidase. Furthermore, the other amino acids mapping to the monomer-monomer interface of ZmGlu1 show high similarity to the corresponding regions of the wheat and rye glucosidases. Thus, the reasons for the weak monomer-monomer interaction within the dimer are unclear at present.

F198A Mutation Reduces the Catalytic Efficiency of TaGlu1a and ScGlu

On replacing the N-terminal 25 residues with the corresponding region of ZmGlu1, the wheat glucosidase (Zm-TaGlu1a or Zm-TaGlu1b) monomer could not make up a hexamer but a dimer (Fig. 4B). The amino acid sequences of the N-terminal region are least conserved among the family GH1 β-D-glucosidases. The dimer of Zm-TaGlu1 easily dissociated into monomers (data not shown), suggesting that the monomer of TaGlu1 interacts very weakly in comparison with the ZmGlu1 or SbDhr1 monomers. These characteristics differ from the monocot family GH1 β-D-glucosidases, where the dimeric forms represent the stable structural units from which the oligo- and multimers are composed. The crystal structure of the TaGlu1b monomer resembles the known structures of the maize and sorghum homologs (Fig. 5C). The two residues in ZmGlu1 (Arg-295 and Asp-342) that were shown to form the intermonomer salt bridges (Czjzek et al., 2001) are conserved in TaGlu1b (Arg-296 and Asp-343). The intramolecular disulfide bond responsible for maintaining the dimeric form of Zm-p60.1 (Zouhar et al., 2001) is also conserved in the wheat glucosidase. Furthermore, the other amino acids mapping to the monomer-monomer interface of ZmGlu1 show high similarity to the corresponding regions of the wheat and rye glucosidases. Thus, the reasons for the weak monomer-monomer interaction within the dimer are unclear at present.

The crystal structure of a β-glucosidase responsible for hydrolysis of Hx-Glc was first solved for maize (Czjzek et al., 2000). The aglycone moiety of DIMBOA-Glc was shown to be stabilized by four aromatic side chains: Trp-378 on one side and three Phe residues (Phe-198, Phe-205, and Phe-466) on the other side (Czjzek et al., 2001; Zouhar et al., 2001). Although Trp-378 and Phe-198 are conserved among the five monocot β-D-glucosidases (glucosidases from wheat, rye, maize, sorghum, and oats), Phe-205 and Phe-466 are substituted by a His and Ser, respectively, in TaGlu1 and His and Gly, respectively, in ScGlu. These amino acid substitutions change the electrostatic and spatial environment of the aglycone binding pocket, although substrate recognition and binding must resemble that of ZmGlu1 because all three enzymes favor Hx-Glc. Assuming the mechanism of substrate recognition is the same for TaGlu1a, ScGlu, and ZmGlu1, the sole aromatic residue (Phe-198) positioned opposite Trp-379 in TaGlu1a and ScGlu is likely to play a significant role in substrate binding. This is supported by the results of the F198A mutants, where the catalytic efficiency ($k_{cat}/K_m$) decreased by more than 95% except for the activity of ScGlu-F198A against pNP-Glc. However, the $k_{cat}$ of ScGlu for DIBOA-Glc and
DIMBOA-Glc was enhanced by this mutation, and the $k_{\text{cat}}$ of TaGlu1a-F198A toward DIMBOA-Glc was about 30% that of the wild-type protein. These results are somewhat different from those for the maize glucosidase, where replacement of Phe-198 by smaller amino acids caused a more drastic reduction in $k_{\text{cat}}$ (Zouhar et al., 2001; Verdoucq et al., 2003). The crystal structure of the ZmGlu1 mutants revealed that the F198V mutation changes the orientation of the side chain of Phe-466, one of the Phe residues constituting the hydrophobic binding pocket, leading to complete loss of activity against pNP-Glc (Verdoucq et al., 2003). In the case of TaGlu1a and ScGlu, the Phe-466 is substituted by Ser and Gly, respectively. Thus, the mutation may alter the environment around the binding pocket to a lesser extent in the wheat and rye glucosidases. Additionally, the Tyr at the entrance to the aglycone binding sites in both glucosidases (Tyr-378) may compensate for the deletion of Phe-198.

**Bulky Aromatic Side Chains at the Entrance of the Substrate Binding Site Play a Significant Role in TaGlu1a**

One of the greatest differences around the aglycone binding pocket between the wheat and rye glucosidases and the maize glucosidase is Tyr-378 in TaGlu1 and ScGlu (equivalent to Pro-377 in ZmGlu1). The bulky side chain of Tyr-378 in TaGlu1a and ScGlu may obstruct access to the binding pocket by the substrate. Indeed, we initially assumed that replacing the Tyr with a small residue might increase the activity of the enzymes. Substitution of the Tyr with Ala caused the $k_{\text{cat}}/K_m$ of ScGlu to increase for DIMBOA-Glc and pNP-Glc by a factor of 4- to 5-fold. However, the efficiency for all substrates was lower for the TaGlu1a-Y378A mutant, suggesting a different role for the Tyr residue in the wheat enzyme. The Ser-464 and Leu-465 residues in TaGlu1a, which are bulkier than the corresponding residues in ScGlu, may require a lid to maintain the substrate in a favorable position. The higher $K_m$ values of the Y378A mutants and the lower $K_m$ values in the Y378F mutants indicate the significance of an aromatic residue positioned at the entrance of the binding pockets. In the case of the maize glucosidase, the lack of an aromatic residue may still give high activity toward DIMBOA-Glc because the binding pockets, comprising four aromatic side chains, can still hold the aromatic aglycone moiety firmly. The lower $k_{\text{cat}}$ values of TaGlu1a- and ScGlu-Y378F mutants compared to those of wild-type proteins may indicate the hydroxy group is necessary for placing the glucosidic bond of the substrate at the optimal position for attack by the catalytic residues. However, a more hydrophobic environment seems to be more favorable for enzyme-substrate binding.

**The Effects of the Mutations at Positions 464, 465, and 471**

This result that the S465L mutation of ScGlu increased the efficiency toward DIMBOA-Glc seems to match with the results of ZmGlu1, where the maize counterpart of the residue is involved in the recognition of the methoxy group in DIMBOA-Glc (Czijzek et al., 2000). However, the single G464S mutation alone could raise the specificity to DIMBOA-Glc. Therefore, amino acid 464 as well as 465 plays an important role in distinguishing DIMBOA-Glc from DIBOA-Glc. The substitutions of Gly-464 and Ser-465 of ScGlu by the TaGlu1 counterparts made the substrate preference of ScGlu resemble that of TaGlu1. However, the $k_{\text{cat}}/K_m$ value toward DIMBOA-Glc was about 30 times larger than that toward DIBOA-Glc in TaGlu1a, although the value against DIMBOA-Glc of ScGlu-G464S/S465L mutant was only about 10 times larger than that against DIBOA-Glc. This would suggest that there are other unknown factors in TaGlu1a that contribute to distinguish DIMBOA-Glc from DIBOA-Glc.

In ZmGlu1, Tyr-473 was shown to form a hydrogen bond with Trp-378, and the mutation of Tyr-473 into Phe resulted in a decrease in efficiency (Verdoucq et al., 2003). The authors suggested that the loss of the hydrogen bond increased the flexibility of Trp-378, resulting in an enhanced catalytic efficiency. Replacing the corresponding residue of TaGlu1a, Phe-471, into Tyr resulted in a decrease in efficiency, which agrees with the results of ZmGlu1. However, the same mutation in ScGlu enhanced the efficiency toward DIMBOA-Glc and pNP-Glc, while it decreased the efficiency toward DIBOA-Glc. It is notable that every ScGlu mutant that showed a higher efficiency toward DIMBOA-Glc than the wild type exhibited an increased value toward pNP-Glc but not toward DIBOA-Glc, although the detailed mechanisms are not known at present.

Because none of the mutations introduced in this study enhanced the efficiency of TaGlu1a, a more precise organization of the substrate binding site is likely to be required to recognize and hydrolyze the substrates in TaGlu1a by comparison with ScGlu. The broader substrate specificity of ScGlu may be due to the wider aglycone binding site compared with those of ZmGlu1 and TaGlu1a. Further structural analyses of TaGlu1 and ScGlu crystals will allow elucidation of the details of substrate recognition and organization of the active hexamers.

**MATERIALS AND METHODS**

**Plant Materials**

Two cultivars of bread wheat (Triticum aestivum), cv Ak and cv CS, and rye (Secale cereale) were grown at 25°C as described previously (Sue et al., 2000a, 2000b).

**Northern-Blot Analysis and Quantitative-PCR**

Total RNA was isolated from wheat shoots (Ak) using RNeasy Plant Mini Kit (QIAGEN). The RNA (15 μg) was separated on a 1.2% agarose MOPS-formaldehyde gel and transferred to a nylon membrane, Hybond N+ (Amersham Biosciences). The membrane was probed with an [α-32P]-labeled cDNA fragment from the 3'-RACE (see supplemental text) as described by Church and Gilbert (1984). The hybridization and washing procedures were...
Expression of the β-d-Glucosidases in Escherichia coli

The full-length cDNAs (Taglu1a-Taglu1c) were used as template for PCR to prepare DNA fragments flanked by Ncol and Xhol recognition sites at the 5′ and 3′ ends, respectively. The sequences of the primers used in the PCR are shown in Supplemental Table I. The primer for Scgulu was designed based on the sequence of the rice glucosidase (GenBank accession no. AF293849) reported by Nikus et al. (2003). The reaction was carried out utilizing KOD polymerase (TOYOBO) with denaturing at 98°C reported by Nikus et al. (2003). The reaction was carried out utilizing KOD polymerase (TOYOBO) with denaturing at 98°C, annealing at 98°C, and polymerization at 74°C. The amplified DNA fragments corresponding to the mature glucosidases were digested by Ncol and Xhol and then cloned into pET21d or pET30a. By introducing the DNA fragments into the Ncol and Xhol sites of pET21d and pET30a, we obtained plasmids for native (without His-tag) and N-terminal His-tagged glucosidases, respectively. The ligation products were transformed into BL21 CondomPlus(DE3)-RIL (competent cells) for coexpression of the glucosidase genes, Taglu1a (or Taglu1c) was cloned into pET21d and Taglu1b into pCDFDuet-1 (Novagen). Both plasmids were then introduced into the BL21-CondomPlus(DE3)-RIL strain. The E. coli was cultured in 50 mL of Luria-Bertani broth supplemented with appropriate antibiotics at 37°C with shaking until the OD600 reached approximately 0.5. Heterologous gene expression was then induced by adding 1 mM isopropyl-β-D-galactoside followed by an overnight culture at 20°C. The cells were pelleted by centrifugation (1,500g for 15 min) and then resuspended in 5 mL of 50 mM HEPES, pH 7.2, containing a protease inhibitor cocktail (Sigma). The cells were disrupted by sonication on ice (several 20-s pulses at a power setting of 100 W). The soluble protein fraction was recovered by collecting the supernatant after centrifugation at 15,000g for 15 min.

The recombinant His-tagged protein was purified by metal chelation chromatography. The resin was washed with a HiTrap Chelating HP column (Amersham Biosciences) equilibrated with 0.02 M phosphate buffer, pH 7.4, containing 0.5 M NaCl and 60 mM imidazole. After washing the column with the same buffer containing 0.5 M NaCl and 60 mM imidazole, the glucosidase was eluted by increasing the concentration of imidazole to 300 mM. The eluate was concentrated by ultracentrifugation and then subjected to gel filtration chromatography on Superdex 200 (Amersham Biosciences) equilibrated with 50 mM HEPES and 130 mM NaCl, pH 7.2. To estimate the molecular mass on the gel filtration column, the following proteins were used as standards: ferritin (440 kDa), human IgG (160 kDa), transferrin (81 kD), ovalbumin (43 kD), and myoglobin (17.6 kD).

Electrophoresis and Activity Staining

The protein profile was analyzed by SDS-PAGE and native-PAGE using an 8% gel as described previously (Sue et al., 2000b). The bands corresponding to β-glucosidase were detected on a native-PAGE gel using a chromogenic substrate, 6-bromo-2-naphthyl-β-D-glucopyranoside, as described previously (Sue et al., 2000b).

Mass Analyses of TagGlula and TagGlub

The purified N-His-tagged TagGlula and TagGlub were concentrated by ultrafiltration, followed by dilution with Milliq water to reduce the salt concentration in the buffer. After several rounds of concentration and dilution, acetonitrile and formic acid were added to give a final concentration of 20% and 0.1%, respectively. The protein concentration was adjusted to 1 mg/mL. The molecular masses of the glucosidases were measured using a Perkin-Elmer-Sciex API-165 (ion-spray voltage 5 kV, orifice voltage 30 V, nebulizer gas N2, curtain gas N2). The theoretical Mw and pI were determined by the Compute pl/Mw program on the ExPaSy server (http://kr.expasy.org/tools/).

Structure Determination

The recombinant β-glucosidase was expressed, purified, and crystallized as described previously (Sue et al., 2005). To obtain the complex of TaGlub and its substrate aglycone, DIBOA, crystals were soaked in the crystallization buffer with 0.5 M DIBMOA and 30% glycerol as a cryoprotectant for 15 min and then cooled in a nitrogen stream at 100 K. The diffraction data set was collected on beamline BL-6A at Photon Factory and processed by the program HKL2000 (Otwinowski and Minor, 1997). The initial model was obtained by the molecular replacement method using the program MOLREP (Vagin and Teplyakov, 1997) and the β-glucosidase molecule from sorghum (Sorghum bicolor; PDB code: 1v03; Verdoucq et al., 2004) as the search model. The crystal belonged to the space group P4232 and contains one monomer in an asymmetric unit. The iterative refinement was performed using the programs CNS (Brünger et al., 1998) and XtalView (McCoy, 1992).

Site-Directed Mutagenesis of His-Tagged TaGlula and ScGlub

Mutated DNA fragments for expression of TaGlula and ScGlub mutants were prepared by PCR-mediated overlap extension. The sequences of the mutagenic PCR primers used in this study are shown in Supplemental Table I. The DNA fragments for ZmTaGlula and ZmTaGlub were amplified by PCR using the primers shown in the Supplemental Table I. The amplified fragments were digested by Ncol and Xhol, introduced into pET30a, expressed in E. coli, and purified by affinity chromatography followed by gel filtration chromatography as described above.

Substrate Preparation and Enzyme Assay

DIBOA-Glc and DIBMOA-Glc were isolated from shoots of 48-h-old rape and maize (Zea mays), respectively, according to methods described previously (Sue et al., 2000b).

Enzyme activity was measured in 100 mM citrate-200 mM phosphate buffer, pH 5.5, at 30°C. The product was quantified by HPLC (elucent, 30% [v/v] methanol containing 0.1% [v/v] acetic acid; column, Inertsil ODs-5 [4.6 × 150 mm; GL Sciences]; temperature, 40°C). The amount of protein and reaction time were carefully chosen so that the product did not exceed 5% to 10% of the residual substrate. Kinetic parameters were determined from several independent experiments. The preliminary Km and kcat values were determined from reactivation rates at various concentrations of substrates ranging from 0.02 mM to 10 mM. Precise kinetic parameters were subsequently obtained by varying the substrate concentration from 20% to 200% of the preliminary Km value. Km and kcat values were calculated by fitting the data to the Michaelis-Menten equation using SigmaPlot 2000 (SYSTAT Software). The protein concentration was measured by the method of Bradford (Bradford, 1976) using bovine serum albumin as a standard.

Modeling of ScGlub

The three-dimensional model of ScGlub was calculated by homology modeling with MODELLER6v2 program (Sali and Blundell, 1993; Marti-Renom et al., 2000) using the structures of TaGlub and ZmGlul (PDB code: 1E56). First, the primary structures of ScGlub, TaGlub, and ZmGlub were aligned using ClustalW, and the result was checked based on the actual (TaGlub and ZmGlub) or predicted (ScGlub) secondary structures. Then, five models were calculated by MODELLER6v2 using the above sequence alignment. Since all structures obtained were almost identical to each other, one model was employed as a predicted structure of ScGlub. The model was evaluated by PROCHECK. Although one residue (Lys-251) was in a disallowed region, it was located on the surface of the protein and far from the substrate binding pocket.

Sequence data from this article can be found in the DDBJ/EMBL/GenBank data libraries under the accession numbers AB100035 (Taglu1a), AB236422 (Taglu1b), and AB236423 (Taglu1c). The atomic coordinates of TaGlub were deposited in the Protein Data Bank under the code 2DGA.
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