Gibberellic Acid-Induced Aleurone Layers Responding to Heat Shock or Tunicamycin Provide Insight into the N-Glycoproteome, Protein Secretion, and Endoplasmic Reticulum Stress

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The growing relevance of plants for the production of recombinant proteins makes understanding the secretory machinery, including the identification of glycosylation sites in secreted proteins, an important goal of plant proteomics. Barley (Hordeum vulgare) aleurone layers maintained in vitro respond to gibberellic acid by secreting an array of proteins and provide a unique system for the analysis of plant protein secretion. Perturbation of protein secretion in gibberellic acid-induced aleurone layers by two independent mechanisms, heat shock and tunicamycin treatment, demonstrated overlapping effects on both the intracellular and secreted proteomes. Proteins in a total of 22 and 178 two-dimensional gel spots changing in intensity in extracellular and intracellular fractions, respectively, were identified by mass spectrometry. Among these are proteins with key roles in protein processing and secretion, such as calreticulin, protein disulfide isomerase, proteasome subunits, and isopentenyl diphosphate isomerase. Sixteen heat shock proteins in 29 spots showed diverse responses to the treatments, with only a minority increasing in response to heat shock. The majority, all of which were small heat shock proteins, decreased in heat-shocked aleurone layers. Additionally, glycopeptide enrichment and N-glycosylation analysis identified 73 glycosylation sites in 65 aleurone layer proteins, with 53 of the glycoproteins found in extracellular fractions and 36 found in intracellular fractions. This represents major progress in characterization of the barley N-glycoproteome, since only four of these sites were previously described. Overall, these findings considerably advance knowledge of the plant protein secretion system in general and emphasize the versatility of the aleurone layer as a model system for studying plant protein secretion.

Plant proteins that are secreted to the apoplast have important functions in signaling, defense, and cell regulation. The classical protein secretory pathway is less characterized in plants than in mammals or yeast but is of growing interest due to the potential of plant systems as hosts for the production of recombinant proteins (Erlendsson et al., 2010; De Wilde et al., 2013). Plant secretomics, therefore, is a rapidly expanding area applied to gain further insight into these processes (Agrawal et al., 2010; Alexandersson et al., 2013). Many secretory proteins contain putative N-glycosylation sites, and the identification and characterization of these sites is an important element in secretomics analysis. However, to date, only a few plant glycoproteomes have been described (Fitchette et al., 2007; Minic et al., 2007; Palmisano et al., 2010; Melo-Braga et al., 2012; Zhang et al., 2012; Thannhauser et al., 2013).

The cereal aleurone layer is of major importance due to its central role in grain germination. Previous proteomic studies have been reported for aleurone layers dissected from mature (Finnie and Svensson, 2003) and germinating (Bonsager et al., 2007) barley (Hordeum vulgare) or developing (Tasleem-Tahir et al., 2011) and mature (Laubin et al., 2008; Jerkovic et al., 2010; Meziani et al., 2012) wheat (Triticum aestivum) grains. In the in vitro culture of isolated aleurone layers was developed by Chrispeels and Varner (1967). Since then, this system has become an excellent tool for the study of germination signaling in response to phytohormones (Bush et al., 1986; Jones and Jacobsen, 1991; Bethke et al., 1997; Ishibashi et al., 2012). More recently, it has been adopted as a unique system for the analysis of plant secretory proteins (Hägglund et al., 2010; Finnie et al., 2011). The addition of GA₃ to the isolated aleurone layers induces the synthesis and secretion of hydrolytic enzymes. In the in vitro system, these accumulate in the incubation buffer, facilitating their identification and characterization using proteomics techniques. Thus, numerous secreted proteins with roles in the hydrolysis of starch, cell wall polysaccharides, and proteins could be identified (Finnie et al., 2011). Furthermore, several of the proteins were also detected in intracellular extracts from...
the same aleurone layers, presumably prior to their release. Many of the proteins appeared in multiple forms on two-dimensional (2D) gels, and often with higher $M_\text{r}$ than expected, suggesting the presence of posttranslational modifications (PTMs; Finnie et al., 2011). Bak-Jensen et al. (2007) and Finnie et al. (2011) observed a highly complex pattern of $\alpha$-amylase-containing spots on 2D gels, which originated from only two $\alpha$-AMYLASE2 (AMY2) and two $\alpha$AMY1 gene products from a total of 10 genes, probably reflecting multiple forms due to PTMs.

In eukaryotic cells, proteins synthesized in the endoplasmic reticulum (ER) must be correctly folded and assembled before continuing in the secretory pathway. Perturbations of redox state, calcium regulation, Glc deprivation, and viral infection can lead to ER stress, triggered by the accumulation of unfolded and misfolded proteins in the ER lumen. This provokes a triple response from the cell, consisting of an up-regulation of chaperones and vesicle trafficking, a down-regulation of genes encoding secretory proteins, and an up-regulation of proteins involved in endoplasmic reticulum-associated protein degradation (ERAD). In plants, the molecular mechanisms underlying ER stress in plants have yet to be fully resolved (Martínnez and Chriseeps, 2003; Nagashima et al., 2011; Moreno et al., 2012).

Tunicamycin (TN), an inhibitor of GlcNAc phosphotransferase, which catalyzes the first step in glycoprotein synthesis, has been used to induce ER stress by causing the accumulation of unfolded proteins in the ER lumen (Noh et al., 2003; Kamauchi et al., 2005; Reis et al., 2011). If unfolded proteins are not removed, the prolonged stress will induce programmed cell death. Links between ER stress and apoptosis have been reported in response to TN treatment in mammalian cells, whereas in plants, this correlation has been suggested, but the pathways of signal transduction remain unknown (Kamauchi et al., 2005; Reis et al., 2011).

In all plant tissues, heat shock (HS) induces the synthesis of a variety of heat shock proteins (HSPs), which are responsible for protein refolding under stress conditions (Craig et al., 1994) and translocation and degradation in a broad array of normal cellular processes (Bond and Schlesinger, 1986; Spiess et al., 1999). In the barley aleurone layer, HS selectively suppresses the synthesis of secretory proteins, including $\alpha$-amylase, due to the selective destabilization of secretory protein mRNA (Belanger et al., 1986; Brodl and Ho, 1991). However, an acclimation effect has been described in aleurone cells after prolonged incubation at warm temperatures, resulting in a resumption of the protein secretory machinery (Shaw and Brodl, 2003). The connection between heat stress response and ER stress has been well established in mammals and yeast, but scarce information is available in plants (Denecke et al., 1995).

Over the last years, the dual role of reactive oxygen species (ROS) has been established in plants: at higher concentrations, ROS act as toxic molecules damaging cellular macromolecules, eventually causing cell death, but at lower concentrations, ROS seem to be necessary for seed germination and seedling growth by controlling the cellular redox status, regulating growth and protecting against pathogens (Bailly, 2004; Bailly et al., 2008). In the barley aleurone layer, GA$_3$ perceived at the plasma membrane induces ROS generation as a by-product from intense lipid metabolism, and the redox regulation of the GA$_3$-induced response has been proposed (Maya-Ampudia and Bernal-Lugo, 2006). This suggests that during the secretory function of the tissue, moderate levels of ROS may be acting as cellular messengers. In aleurone cells, ROS, especially hydrogen peroxide (H$_2$O$_2$), are involved in the process of programmed cell death, but the molecular mechanisms remain unclear (Bethke and Jones, 2001; Ishibashi et al., 2012).

Until now, none of the protein components of the ER stress pathways have been identified in barley; also, little is known about the glycosylation of barley proteins. In this work, numerous $N$-glycosylation sites are identified, and the effect of perturbing $N$-glycosylation and the secretory pathway by TN and HS treatments is analyzed in GA$_3$-induced barley aleurone layers.

RESULTS

The Effect of TN and HS on $\alpha$-Amylase Production in GA$_3$-Induced Aleurone Layers

Since $\alpha$-amylase is the most prominent secreted enzyme produced by aleurone cells in response to GA$_3$, a perturbation of the secretory machinery by TN or HS may affect the amount of this enzyme. Western blotting was used to detect $\alpha$-amylase in the extracellular and intracellular fractions from aleurone layers incubated for 24 h. Untreated, GA$_3$-induced, and GA$_3$-induced aleurone layers in combination with HS or TN treatment were compared (Fig. 1). A decrease in $\alpha$-amylase could be observed, both in the extracellular (Fig. 1A) and intracellular (Fig. 1B) fractions, when GA$_3$-induced cells were treated with 5 or 20 $\mu$g mL$^{-1}$ TN. Higher TN concentrations did not elicit a further decrease in the amount of $\alpha$-amylase (data not shown).

For HS treatment, a temperature of 40°C was applied for the final 4 h of the 24-h incubation period. The HS treatment caused a decrease in $\alpha$-amylase in the intracellular fraction (Fig. 1B) and in $\alpha$-amylase secreted during the final 4 h when compared with GA$_3$-induced aleurone cells (Fig. 1A). Longer incubation periods at 40°C had no apparent effect on the amount of $\alpha$-amylase (data not shown), most likely due to the acclimation response reported previously (Shaw and Brodl, 2003).

GA$_3$-Induced Cell Death Correlates with Enhanced H$_2$O$_2$ and Lipid Peroxidation

Both H$_2$O$_2$ content and thiobarbituric acid-reactive substances (TBARS) indicating lipid peroxidation were increased in extracts from GA$_3$-induced aleurone layers in comparison with untreated samples (Fig. 2A). Treatment of GA$_3$-induced aleurone layers with 5 $\mu$g mL$^{-1}$ TN or with HS did not result in further increases, but
20 μg mL⁻¹ TN significantly enhanced the level of lipid peroxidation. In order to examine whether TN and HS could affect the viability of barley cells, cell death was monitored and quantified in intact aleurone layers by simultaneous staining of live and dead cells with fluorescent probes (Fig. 2B). The cell death quantitation reflected a correlation with the endogenous H₂O₂ and lipid peroxidation measurements. GA₃ treatment of barley aleurone cells resulted in increased cell death, while untreated aleurones maintained higher cell viability. Hence, after 24 h of incubation, 45% of GA₃-induced cells were dead, whereas 81% of untreated cells remained alive (Fig. 2B). GA₃ + 20 μg mL⁻¹ TN did not result in increased cell death, whereas cell death was enhanced in GA₃ + 20 μg mL⁻¹ TN samples (Fig. 2B). Treatments of GA₃-induced aleurone layers with 5 μg mL⁻¹ TN and 4 h of HS were selected for comparative proteome analysis, since these affected protein secretion without provoking massive oxidative damage and cell death.

Fluorescent Glycoprotein Staining and Protein Identification

To visualize glycoproteins and assess the effect of TN treatment on protein glycosylation, extracellular proteins from GA₃-induced and GA₃ + 5 μg mL⁻¹ TN-treated aleurone layers were resolved by 2D SDS-PAGE, and the glycoproteins were detected by Pro-Q Emerald staining. As expected, TN greatly reduced the glycosylation of extracellular proteins (Fig. 3A). The fact that some highly abundant proteins did not react with the Pro-Q Emerald stain whereas some low-abundance proteins reacted strongly supports the specificity of Pro-Q Emerald staining for glycoproteins. To reduce the risk of false positives, of the 53 fluorescent spots visible in all three GA₃-treated replicates, only the 23 intense spots missing after TN treatment (numbered spots in Fig. 3B) were analyzed by mass spectrometry. Proteins were identified in 14 spots (Supplemental Table S1). Sequence analysis (http://www.cbs.dtu.dk/services/NetNGlyc/) confirmed the presence of at least one putative N-glycosylation site in each identified protein (data not shown).

Proteome Analysis of the Extracellular Fraction from in Vitro Incubated Aleurone Layers

2D gel electrophoresis (pH 3→10) was used to profile the extracellular protein fraction from the four aleurone layer treatments. Clear differences were apparent between 2D gel electrophoresis patterns of untreated and GA₃-induced aleurone layers as well as between GA₃-induced aleurone layers and each stress treatment (Fig. 4A). In total, the average volumes of 35 spots varied by at least 1.5-fold among the four treatments. These spots were chosen for further analysis, and 22 were identified...
by mass spectrometry (Fig. 4B; Supplemental Table S2). As expected, most of them were α-amylases and proteases (Fig. 4C). The spots formed three clusters according to their appearance profiles (Fig. 5). Protein spots in cluster A (13 spots) increased in response to GA3: 11 contained α-amylases (AMY1 and AMY2) and one contained a Cys proteinase (EP-B1). Moreover, four spots decreased in GA3 + TN samples (cluster B). Here, one AMY2 spot migrated more slowly than the other identified AMY1 and AMY2 spots (Supplemental Table S2), indicating that this upper spot may be glycosylated. Finally, five spots decreased in intensity in both treatments (cluster C). Cluster C contains four protease spots (two Cys proteinase and two cathepsin) with higher experimental Mr than Cys proteinase spots in clusters A and B, again suggesting the presence of PTMs (Supplemental Table S2).

Proteome Analysis of Aleurone Layer Intracellular Extracts

Intracellular water-soluble proteins from the four samples described above were also analyzed using 2D gel electrophoresis (Fig. 6A) and mass spectrometry. The image analysis showed 250 protein spots varying among the four treatments (Fig. 6B). Of these, 178 were identified (Fig. 6B; Supplemental Table S3) and assigned to seven functional categories (Fig. 6C). Among the identified proteins, 55% (98 protein spots) are involved in primary metabolism; the remaining proteins were functionally assigned as chaperones (29), polysaccharide hydrolases (11), detoxification enzymes (12), signaling proteins (nine), and defense proteins (four), and a considerable number of identified spots (16) were proteins of unknown function (Fig. 6C). Principal component analysis (PCA) performed on these differentially changed spots (Fig. 7A) indicated the presence of six clusters based on similarity of expression profiles (Fig. 7B). Proteins in cluster A (decreased in GA3 and GA3 + TN aleurone samples and increased in GA3 + HS samples) and in cluster B (increased in GA3 + HS samples) were mainly proteins involved in primary metabolism, such as glycolysis, citric acid cycle, and amino acid biosynthesis. Remarkably, 11 out of 22 spots in cluster C with lower abundance in GA3 + HS samples contain small HSPs. In cluster D, the spots from the three samples incubated
with GA3 decreased in volume when compared with untreated samples; it is also very interesting that 38% of the identifications in cluster D were proteins of unknown function. Clusters C and E grouped protein spots whose profiles dramatically decreased in HS samples when compared with GA3- and GA3 + TN-treated samples. Proteins in 28 spots in cluster E belong to primary metabolism, indicating heat-induced protein degradation; remarkably, three AMY1 and three AMY2 spots were assigned to cluster E, suggesting selective destabilization of secretory protein mRNA by heat stress. Finally, spots showing increases both in GA3 + TN and GA3 + HS samples were grouped in the small cluster F, including several late embryogenesis abundant

Figure 4. Extracellular protein profiles of barley aleurone layer extracts. A, One representative colloidal Coomassie blue-stained gel covering the pH range 3 to 10 is shown for each treatment. Molecular size markers are indicated at left. U, Untreated. B, Closeup view of the reference gel showing 22 identified spots. C, Functional categories of the plant proteins identified. (For details about protein identifications, see Supplemental Table S2).
proteins and protein disulfide isomerase (PDI). Furthermore, among the six clusters there were 19 protein spots suspected to be protein fragments due to a clear discordance between theoretical and experimental pI and Mr (Supplemental Table S3). In accordance with this, their sequence coverages obtained by Mascot search were confined to the N- or C-terminal parts of the protein. Curiously, 17 of these 19 protein fragments belong to cluster E.

Identification of N-Glycosylated Proteins from Complex Protein Samples

An N-glycoproteome analysis was performed in both intracellular and extracellular fractions from aleurone layers in order to provide the first overview of glycosylation sites in barley proteomes. Enrichment of glycopeptides from trypsin-digested protein extracts was carried out by hydrophilic interaction liquid chromatography (HILIC), followed by deglycosylation using PNGase A in the presence of H$_2^{18}$O. During the deglycosylation reaction, the formerly glycosylated Asn residues undergo deamidation and $^{18}$O is incorporated into the carboxylate groups of the resulting Asp side chains. Deglycosylated Asn residues can thus be distinguished from residues that were deamidated prior to the PNGase A reaction. Analysis by liquid chromatography (LC)-tandem mass spectrometry (MS/MS) on a Q-Exactive Orbitrap enabled the determination of 73 independent glycopeptides in 65 glycoproteins (Table I; Supplemental Table S4). Of these, 57 glycoproteins were found in the extracellular fraction, whereas 36 glycoproteins were identified in the intracellular samples. A total of 28 proteins were found in both fractions. All the glycoproteins were predicted to follow the secretory pathway according to the signal peptide cleavage site prediction (Supplemental Table S4).

DISCUSSION

By the application of two stresses that affect protein secretion through independent mechanisms, new
Figure 6. Intracellular water-soluble protein profiles of barley aleurone layer extracts. A, One representative colloidal Coomassie blue-stained gel covering the pH range 3 to 10 is shown for each treatment. Molecular size markers are indicated at left. U, Untreated. B, Closeup view of the reference gel showing 178 identified spots. C, Functional categories of the plant proteins identified. (For details about protein identifications, see Supplemental Table S3).
information about the plant protein secretion system in GA$_3$-induced aleurone layers has been obtained.

A clear effect of GA$_3$ on H$_2$O$_2$ production and cell death was confirmed (Bethke and Jones, 2001; Ishibashi et al., 2012). Moreover, lipid peroxidation, as indicative of cell damage in response to several stresses (Bailly, 2004; Barba-Espín et al., 2011), was enhanced by GA$_3$ (Fig. 2A). A high TN concentration (20 µg mL$^{-1}$) provoked further H$_2$O$_2$ accumulation and cell death, which is probably due to the oxidative damage enhanced in these samples. The inhibition of protein folding and blocking of secretion provoked by high TN concentrations seemed to be incompatible with cellular function, whereas the response triggered by lower TN concentrations may still allow recovery of the ER function.

Overall, GA$_3$ induced protein secretion from aleurone layers as shown (Finnie et al., 2011). Half of the observed spots decreased in abundance under one or both of the stress treatments. Most of these contained proteases, whereas the majority of the α-amylase spots remained at similar levels to unstressed (GA$_3$-induced) aleurone layers (Figs. 4 and 5; Supplemental Table S2). Cys proteinases are the most abundant proteases secreted by barley aleurone layers in response to GA$_3$ (Zhang and Jones, 1995; Martinez and Diaz, 2008), and the synthesis of several family members remains high during the first 36 h of GA$_3$ incubation (Koehler and Ho, 1990). By contrast, α-amylase secretion peaks much earlier during the first 24 h of incubation (Bethke et al., 1997). Thus, the difference in the responses of proteases and α-amylase to the treatments could be due to more significant secretion of Cys proteinases than α-amylases during the 4-h period of HS.

In the intracellular proteomes, 178 spots changed abundances and were assigned to six clusters (Figs. 6 and 7). The overall effect of TN and HS treatments on the intracellular spot patterns reflects a differential protein response; HS provoked major changes in the protein patterns when compared with treatments with GA$_3$ alone and TN, as expected due to the suppression of secretory protein synthesis (Brodl and Ho, 1991; Spiess et al., 1999) and the more selective mechanism of the TN effect on the ER lumen. Considering the short incubation time at 40°C, the major changes occurring in HS samples suggest changes in protein turnover. In fact, the most dramatic variations between GA$_3$ and stress treatments correspond to spots either decreasing or increasing in HS samples (Fig. 7; Supplemental Figure 7. PCA and clustering of intracellular water-soluble protein spots. A, PCA was performed on the 178 identified spots. Biological replicates are grouped by circles, and spots are indicated by numbers. B, Expression profiles of protein spots from PCA analysis were grouped in six clusters. The normalized volume for each spot is expressed relative to a reference gel (GA$_3$), in which all spot volumes are by default set to 1. Corresponding functional classifications are indicated: CH, Chaperone; DE, defense; DT, detoxification enzymes; PH, polysaccharide hydrolase; PM, primary metabolism; SN, signaling; UF, unknown function. U, Untreated. (For details about protein identifications, see Supplemental Table S3).
Table S3). Complementary expression profiles are observed for clusters A and E (Fig. 7), reflected by several full-length proteins in cluster A (aconitate hydratase, HSP70, enolase, aldose reductase, 6-phosphogluconate dehydrogenase, phosphoenolpyruvate carboxylase, and malic enzyme) being present in cluster E as probable fragments.

The aleurone layer contains large reserves of oil, and GA3 stimulates both fatty acid β-oxidation and the glyoxylate cycle for the production of Suc from storage lipids and to produce carbon skeletons for the production of secreted hydrolases (Doig et al., 1975; Eastmond and Jones, 2005). The glyoxylate cycle enzymes isocitrate lyase and aconitate hydratase were identified in 12 and eight spots, respectively, with a number of these containing fragments (Supplemental Table S3). Those migrating according to their predicted M, pl all decreased in intensity in GA3 + TN samples, as did a predicted acyl-CoA oxidase identified in one spot (Supplemental Table S3). This suggests a down-regulation of β-oxidation and the glyoxylate cycle coupled to the decrease in production of secretory proteins.

Table 1. A total of 65 N-glycoproteins was identified from extracellular and intracellular complex samples

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In agreement with previous work (Finnie et al., 2011), several secretory proteins (AMY1 and AMY2, α-N-arabinofuranosidase A, β-xylosidase, and Ser carboxypeptidase) could be identified in intracellular aleurone layer extracts (Supplemental Table S3). These intracellular forms are presumably en route in the protein secretion pathway. Spots containing AMY1 and AMY2, α-N-arabinofuranosidase A, and β-xylosidase were decreased in the intracellular extracts subjected to TN or HS treatments. This decrease was not seen in the extracellular protein fractions (Fig. 4; Supplemental Table S2), probably reflecting that the secretory proteins are transiently present within the aleurone layers, while the extracellular fraction contains proteins accumulating over the entire incubation period. For this reason, it was not possible to observe a decrease in α-amylase after 4 h of HS by western blotting of extracellular proteins accumulated over 24 h (data not shown), although the decrease was apparent both in the intracellular fraction (Fig. 1B) and in the extracellular fraction collected solely during the 4-h period of HS (Fig. 1A).
Many of the extracellular and intracellular proteins (Supplemental Tables S2 and S3) migrated on gels with higher M, than predicted, suggesting the presence of PTMs. These proteins were confirmed to have a leader peptide-processing site and at least one putative N-glycosylation site (data not shown). Information about the PTMs of barley α-amylases is lacking, but extensive modifications are suggested by the complex pattern of spots detected in both supernatant and intracellular fractions (Bak-Jensen et al., 2007; Finnie et al., 2011).

According to Swiss-Prot database predictions, more than 50% of eukaryotic proteins are glycosylated (Apweiler et al., 1999), the majority of them being related to the secretory system. Therefore, a large number of proteins are potential targets for TN action. Many studies have characterized a high number of glycoproteins from bacterial (Nostaft and Szymanski, 2010) and animal (Bunkenborg et al., 2004; Wollscheid et al., 2009; Zielinska et al., 2010) species. This work represents a major contribution toward mapping the barley glycoproteome, assigning 73 N-glycosylation sites in 65 proteins (Table I; Supplemental Table S4), several of which were found by 2D-based analysis either to increase (PDI and calreticulin) or decrease (β-xylosidase, α-N-arabinofuranosidase A, and Ser carboxypeptidase) in response to TN or HS treatments (Fig. 7; Supplemental Table S3). According to Uni-Prot database annotations, from the 65 assigned glycoproteins, only β-xylosidase, Ser carboxypeptidase, xylanase inhibitor, and oxalate oxidase were previously confirmed to be N-glycosylated in barley.

TN treatment of GA3-induced aleurone layers resulted in a decrease in Pro-Q Emerald-stained glycoproteins in the extracellular fraction, including cathepsin B, Cys proteinase C1A, two α-amylase (AMY2), two purple acid phosphatase, two subtilisin-like Ser protease, and six β-glucosidase spots (Fig. 3; Supplemental Table S1). Of these, a corresponding glycopeptide was identified in a Cys proteinase, purple acid phosphatase, subtilisin-like Ser protease, and a β-glucosidase (Supplemental Table S4).

To date, no α-amylase form has been confirmed to be glycosylated. The barley seed complex 2D spot pattern obtained from a limited number of gene products (Bak-Jensen et al., 2007; Finnie et al., 2011) probably reflects PTMs. Here, we show by specific fluorescent staining that two AMY2 spots contain glycoproteins. In agreement with this, both forms migrated with an apparent molecular mass 2 kD larger than the theoretical value. Moreover, in GA3-induced aleurone layers treated with TN, the corresponding AMY2 spots with elevated M, decreased in intensity for both extracellular and intracellular fractions, representing the most significant intensity change in response to TN. By contrast, AMY2 forms migrating according to their predicted M, were not affected by TN. A single N-glycosylation site is predicted in this AMY2 gene product; however, a corresponding glycopeptide was not identified. This may be due to the high hydrophilicity of this particular glycopeptide, hindering its retention on the reverse-phase column prior to mass spectrometry analysis. The glycosylated AMY2 forms identified here constitute a minor proportion of the total α-amylase produced by the isolated aleurone layers (Fig. 3) or in germinating grains (Bak-Jensen et al., 2007). Interestingly, the presence of glycosylated AMY2 varies among barley cultivars, since the AMY2 form with elevated M, was only seen in some cultivars (Bak-Jensen et al., 2007). It remains to be determined whether the glycosylation affects the thermostability or kinetic properties of barley AMY2, as suggested for rice (Oryza sativa) Amy1A (Terashima et al., 1994).

Plant cells seem to have a common strategy for overcoming ER stress through (1) attenuation of genes encoding secretory proteins, (2) enhancement of protein-folding activity by chaperones, such as the ER-localized calreticulin or PDI, and (3) degradation of unfolded proteins by means of the ERAD system (Travers et al., 2000; Martínez and Chrispeels, 2003; Kamauchi et al., 2005). This study reveals components and evidence of this triple response. Five major HSP families are conserved among species (Wang et al., 2004): HSP60, HSP70, HSP90, HSP100, and small HSP (12–40 kD). Marked differences in proteins with chaperone functions were observed between TN and HS samples, suggesting different pathways in the induction of HSPs and the ER stress response (Johnston et al., 2007). Whereas TN treatment increased nine HSP spots (17, 26, and 70 kD), HS induced an increase in six HSP spots (17, 60, and 70 kD) but a decrease in 17 HSP spots (mainly small HSP). Thermoprotection in the aleurone has been reported to correlate more with enhanced levels of fatty acid saturation in membrane phospholipids than with HSP expression, since HSP70 synthesis was only slightly induced by 3 h of incubation at 40°C (Shaw and Brodl, 2003). Our results suggest an overall decrease of small HSP and an increase in certain intermediate- and high-M, chaperones in HS samples; these observations indicate the differential HSP expression under HS and highlight the secondary role of HSPs in acquiring thermotolerance in barley aleurone cells. Moreover, five spots corresponding to HSP70 fragments decreased intensity in HS samples. In fact, some other metabolic protein spots judged to be the result of proteolytic processing were decreased after HS treatment, suggesting changes in protein turnover (Fig. 7; Supplemental Table S3).

Two important ER stress-induced chaperones were found to increase in response to both TN and HS treatments: calreticulin and PDI. Family members of these ER-resident multifunctional proteins are known to be specifically activated by INOSITOL-REQUIRING ENZYME1/basic Leucine Zipper60 (IRE1/bZIP60) transcription factor, which is a mediator of the ER stress response (Ye et al., 2011; Moreno et al., 2012). Calreticulin was identified in two spots with the same pI on 2D gels but differing in molecular mass by about 2 kD, suggesting that the protein in the slower migrating
spot may be N-glycosylated and that in the faster migrating spot may be a nonglycosylated form. In agreement with this hypothesis, TN caused a specific increase in intensity of the lower spot, which was the most pronounced accumulation observed in response to TN, whereas the intensity of the upper spot decreased in abundance. By contrast, HS slightly reduced the level of the lower calreticulin spot and increased the upper spot (Supplemental Table S3). Thus, HS and TN have contrasting effects on the two calreticulin-containing spots. Moreover, PDI was identified in one spot increasing upon both TN and HS treatments (Supplemental Table S3). Glycosylation of both calreticulin and PDI was confirmed by identification of the predicted glycosylation site (Supplemental Table S4). Interestingly, the glycopeptides were identified in the extracellular fraction. The significance of this is unclear; however, calreticulin is known to have multiple cellular locations and in mammalian cells has been identified at the cell surface (Gardai et al., 2005; Gold et al., 2010). A further approach to gain insight on the biological role of calreticulin glycosylation is needed, but an effect on its interaction with other proteins and subcellular localization is expected. Two proteasome subunit spots were also induced, one of them upon TN treatment and the other after HS, both likely associated with the ERAD system.

Two 1-Cys peroxiredoxin spots differing in pI were identified (Supplemental Table S3). Besides its antioxidant role during barley germination (Stacy et al., 1999), a chaperone activity was demonstrated for 1-Cys peroxiredoxin from Chinese cabbage (Brassica campestris ssp. pekinensis) seeds (Kim et al., 2011). 1-Cys peroxiredoxins are susceptible to overoxidation of the active-site Cys, which can be observed in 2D gel electrophoresis, since this causes a pI shift of the protein (Pulido et al., 2009). Chaperone activity was increased in the oxidized form, whereas the reduced form had greater peroxidase activity (Kim et al., 2011). In our experiments, these reduced and oxidized forms likely correspond to the high- and low-pI spots, respectively. HS and to a lesser extent TN treatment caused an increase in the reduced form and a decrease in the oxidized form, which might indicate a decrease in its chaperone activity and an enhanced antioxidant activity.

Cluster F (Supplemental Table S3) contained proteins in addition to PDI that increased in response to both stress treatments and, therefore, are strong candidates for having roles in ER stress mechanisms. Some of these are known to respond to various abiotic stresses, such as Glc and ribitol dehydrogenase (Witzel et al., 2010) and glyoxalase (Wu et al., 2013). Interestingly, a Rib-phosphate pyrophosphokinase (phosphoribosyl pyrophosphate synthase) was identified, which synthesizes phosphoribosyl pyrophosphate, an important intermediate in purine and pyrimidine biosynthesis (Zrenner et al., 2006) The increase in the amount of this protein could suggest a need for DNA repair. A putative isopenetyl diphosphate isomerase was also identified in this cluster (Supplemental Table S3).

This enzyme belongs to the mevalonate pathway and is required for the synthesis of the polyisoprenoid lipid carrier for the glycans used in N-glycosylation (Jones et al., 2009). Therefore, a compensatory mechanism to restore the folding capacity of the ER might be suggested, although further approaches will be required to analyze the mevalonate pathway activity under stress conditions.

Notably, numerous proteins induced by TN treatment, including small HSPs and calreticulin (Supplemental Table S3), were also slightly up-regulated in GA<sub>3</sub>-induced compared with untreated aleurone layers. This was reflected in the PCA, where grouping of the GA<sub>3</sub>-induced and GA<sub>3</sub> + TN-treated aleurone layers suggested similar changes in comparison with the untreated samples, in contrast to the GA<sub>3</sub> + HS-treated aleurone layers (Fig. 7A). This suggests that the GA<sub>3</sub>-induced aleurone layers may display early symptoms of ER stress due to the heavy load on the protein secretory machinery (Fig. 8). The experimental conditions used here were carefully chosen to avoid the induction of cell death due to ER stress but to inflict a subtle pressure on the protein secretory system. Further investigations along these lines of evidence will explore possible links between GA<sub>3</sub>-induced cell death and ER stress in barley aleurone layers.

CONCLUSION

Overall, the data presented here provide new insights into the proteomes of in vitro maintained aleurone layers. (1) A large number of proteins have been identified in the intracellular water-soluble fraction, many of which had previously not been identified in barley aleurone layers. (2) We provide, to our knowledge, the first comparative study of the distinct and overlapping effects of TN and HS on a plant proteome and show that this strategy identifies key components of the protein secretory and quality-control machinery and also identifies proteins with likely functions in ER stress responses. In summary, HS entails a major

![Figure B](https://www.plantphysiol.org)
degradation of the water-soluble protein fraction when compared with TN-treated samples, whereas the TN effect was more selective. In particular, distinct subsets of HSPs respond divergently to HS. Common components of the ER stress response induced by the TN and HS treatments include HSPs, calreticulins, enzymes of nucleotide metabolism, and isopentenyl diphosphate isomerase, and a reduction is observed in polysaccharide hydrolases and proteases in both extracellular and intracellular fractions. (3) The assignment of N-glycosylation sites represents a major advance in mapping the barley glycoproteome, determining the largest number of N-glycoproteins in this species to date. This approach provides targets for further proteomic experiments and the characterization of specific protein patterns.

In conclusion, our work emphasizes new features of the barley aleurone layer, expanding its use as a suitable model for the analysis of the plant protein secretion system and the nature of plant PTMs.

MATERIALS AND METHODS

Plant Material, Experimental Design, and Sample Preparation

Barley grains (Hordeum vulgare ‘Himalaya’, 2003 harvest), representing a pool of individuals, were purchased from Washington State University. These were used as the starting point from which replicate samples were prepared. Aleurone layers were isolated as described previously (Hynek et al., 2006). For each treatment, 100-mg aleurone layers were incubated in 2 mL of a baseline culture medium (20 mM sodium succinate and 20 mM CaCl2, pH 4.2) and incubated at room temperature with continuous gentle shaking for 24 h. When 5 μM GA3 was added to the baseline culture, samples are referred to as GA3-induced aleurone layers. Two additional treatments were applied to aleurone layers incubated in baseline culture + 5 μM GA3: either TN (added from a stock in methanol) or HS (40°C during the last 4 h of incubation).

Aleurone layers and incubation buffers containing the extracellular proteins from the above samples (untreated, GA3-induced, GA3 + HS, and GA3 + TN) were harvested at 24 h, and samples were prepared as follows.

Intracellular Protein Fraction

Aleurone layers were washed four times with 2 mL of the baseline medium, frozen in liquid nitrogen, and stored at −80°C until use. Proteins were extracted from 100-mg aleurone layers in 1 mL of buffer (5 mM Tris-HCl and 1 mM CaCl2, pH 7.5; Østergaard et al., 2002) with the protease inhibitor cocktail Complete (Roche) for 30 min in a cold room. Insoluble material was pelleted by centrifugation (10 min at 15,000 rpm, 4°C), and supernatants containing solubilized proteins were stored at −80°C until use.

Extracellular Protein Fraction

The spent incubation medium from the aleurone layers was centrifuged to remove insoluble material (5 min at 13,000 rpm, 4°C), and then supernatants containing the extracellular proteins were frozen at −80°C until use. At least three biological replicates were used for each analysis. In each one, the whole procedure, starting from the selection of the dry seeds, was repeated.

Protein Quantification

Protein concentrations in both intracellular and extracellular protein fractions were determined using the assay of Popov et al. (1975) with bovine serum albumin as a standard.

Determination of Cell Viability and Death

The percentages of live and dead cells were determined by double staining with fluorescein diacetate (2 μg mL−1 in 20 mM CaCl2) for 15 min, followed by N-(3-Triethylammoniumpropyl)-4-(6-4-(diethylamino)phenyl)hexatrienyl) pyridinium dibromide (20 μM in 20 mM CaCl2) for 2 min as described (Fath et al., 2001; Wu et al., 2011) with minor modifications. Aleurone layers were observed with a fluorescence microscope (Nikon Eclipse E1000; Nikon Instruments) using a 20× objective, and images were captured using a digital camera. Randomly selected fields from three different aleurone layers per treatment (biological replicates) were counted to determine the percentage of live cells.

Determination of Endogenous H2O2 and Lipid Peroxidation (TBARS)

The measurement of H2O2 in the aleurone layers was based on the peroxide-mediated oxidation of Fe3+, followed by the reaction of Fe3+ with xylene orange (Bellincampi et al., 2000). Absorbance values were calibrated to a standard curve generated with known concentrations of H2O2.

The extent of lipid peroxidation was assayed by determining the concentration of TBARS as described (Hernández and Almansa, 2002). Four biological replicates of 20 aleurone layers per treatment were used for each analysis. Volumes of the samples and reactants were adapted for measuring the absorbance on 96-well microplates. The statistical significance of the results was analyzed according to Tukey’s test (P < 0.05) using the XLSTAT software for Mac (Addinsoft).

Western Blotting for α-Amylase Detection

Either 5 μg of protein extracted from the aleurone layer (intracellular fraction) or proteins contained in 20 μL of the extracellular fraction was separated on 4% to 12% Bis-Tris NuPAGE gels (Invitrogen) followed by electroblotting to a nitrocellulose membrane (Hybond-ECL; Amersham, GE Healthcare) before being probed with rabbit polyclonal antibodies raised against barley α-amylase (Søgaard and Svensson, 1990). Secondary goat anti-rabbit IgG antibodies conjugated to alkaline phosphatase and 5-bromo-4-chloro-3-indolyl phosphate/nitroblue tetrazolium western-blotting reagent (Sigma Fast; Sigma-Aldrich) were used to detect binding. Three biological replicates were performed. For each one, a gel was run in parallel and stained with silver nitrate (Hekeshoven and Dernick, 1988). For analysis of the effect of HS on the extracellular α-amylase, the incubation buffer was replaced at the onset of the HS, allowing harvest of the proteins released only during the final 4 h of incubation. To enable sufficient protein to be loaded on the gel, it was necessary to concentrate these protein samples by precipitation (4 volumes of acetone at −20°C overnight).

2D Gel Electrophoresis and Staining

Gel for Mass Spectrometry Analysis

Both the secreted proteins present in 2 mL of incubation buffer and proteins from aleurone layer extractions were desalted on NAP-5 columns (GE Healthcare) following the manufacturer’s instructions. Aliquots containing 140 μg of proteins were precipitated with 4 volumes of acetone and applied to 18-cm pH 3 to 10 Linear Immobilized pH Gradient (IPG) strips (Immobiline DryStrip; GE Healthcare) for mass spectrometry analysis. After scanning, gel images were analyzed using the Progenesis SameSpots (version 4.5.325.33621; Nonlinear Dynamics). Statistical comparison of the spot intensity among the four treatments, based on one-way ANOVA (P < 0.05) and a fold change greater than 1.5 (calculated as the difference between the lowest and the highest mean normalized spot volumes), was verified using false discovery rate analysis (q < 0.05).

Gel Glycoprotein Staining and Fluorescence Visualization

Secreted proteins from GA3 and GA3 +5 μg mL−1 TN samples (100 μg) were precipitated and run on 11-cm pH 3 to 10 Linear IPG strips as described above. The IPG strips were then trimmed to 7 cm, covering pH range 4 to 8.5, and the second-dimension SDS-PAGE (NuPAGE Novex 4-12% Bis-Tris ZOOM Gel; Invitrogen) was performed in the XCell SureLock Mini-Cell...
system (Invitrogen). Gels were stained with the Pro-Q Emerald 300 Glycoprotein Gel Kit (Molecular Probes) according to the manufacturer’s instructions. Three biological replicates were performed for each of the two treatments. The fluorescent spots were visualized using the UVP BioSpectrum Image System (AH Diagnostics) with transillumination at 302 nm. Images of gels were obtained using a 485- to 655-nm filter. After imaging, the gels were stained with colloidal Coomassie Brilliant Blue.

In-Gel Digestion and Protein Identification by Matrix-Assisted Laser-Desorption Ionization Time of Flight/Time of Flight

Spots of interest were excised from 2D SDS-PAGE gels and digested as described by Shevchenko et al. (1996). After soaking with trypsin (modified porcine trypsin; Promega), the gel pieces were covered with 70 µL of 10 mM (NH₄)₂HCO₃ and incubated at 37°C overnight. The supernatant containing tryptic peptides was transferred to a clean tube, and 2 µL was placed onto a matrix-assisted laser-desorption ionization anchor target. The sample was dried at room temperature and covered with 1 µL of 5 µg µL⁻¹ α-cyano-hydroxysynthetic acid in 70% (v/v) acetonitrile (ACN) and 0.1% (v/v) trifluoroacetic acid, dried again, and washed with 2 µL of 0.5% (v/v) trifluoroacetic acid. An Ultraflex II matrix-assisted laser-desorption ionization time of flight/time of flight mass spectrometer (Bruker-Daltonics) was used in positive ion reflector mode, and spectra were analyzed using FlexAnalysis software (Bruker-Daltonics). Internal calibration was carried out with trypsin autolysis products (mass-to-charge ratios 842.51 and 2211.10).

The peptide masses and sequences obtained were matched to proteins with an in-house Mascot server (version 2.2.04), searching in the Munich Information Center for Protein Sequences barley genome database containing high-confidence gene models (ftp://ftpmpis.helmholtz-muenchen.de/plants/barley/public_data/). Search parameters were as follows: allowed modifications, oxidation of Met (variable) and carbamidomethylation of Cys (static); monoisotopic mass tolerance, 40 ppm; allowed missed cleavages, one.

Positive protein identifications were considered when one of the following requirements was met: three independent peptides, where one of them was confirmed by MS/MS fragment ion fingerprinting; or two independent peptides, both confirmed by MS/MS. For details, see Supplemental Tables S1 to S3.

Identification of N-Glycosylated Proteins from Complex Samples and Assignment of Their Glycosylation Sites

In-Solution Digestion and HILIC

Protein samples from both intracellular and extracellular fractions were digested in solution as described (Haggland et al., 2004) and applied to microcolumns containing a combination of Zwitterion-HILIC resin and cotton. A small piece of cotton wool (approximately 200 µg) taken from a cotton wool pad was pushed into a 20-µL GELoader tip (Eppendorf), and ZIC-HILIC resin (Sequant; particle size, 10 µm) was packed on top of the cotton wool. The microcolumns were equilibrated with 40 µL of 80% (v/v) ACN and 0.5% (v/v) formic acid (FA). In-solution trypsin-digested protein samples (20 µg) were dried (SPD 1010 Speed Vac; Thermo Scientific), redissolved in 40 µL of 80% (v/v) ACN and 0.5% (v/v) FA, and applied onto the equilibrated HILIC microcolumns. The column was washed twice with 80% (v/v) ACN and 0.5% (v/v) FA, followed by elution of bound peptides in 0.5% (v/v) FA.

Endoglycosidase Digestion and Protein Identification by Q-Exactive LC-MS/MS

Deglycosylation of the eluted peptides was performed in 100 mM ammonium acetate in H₃BO₃-water for 24 h at 37°C with 0.1 unit of N-glycosidase A (PNGase A; Roche). To minimize 18O isotope labeling at peptide C termini, the deglycosylated samples were dried, redissolved in 100 mM ammonium acetate prepared in H₃BO₃-water, and incubated for 24 h at 37°C in the presence of 0.2 µg µL⁻¹ sequencing-grade modified trypsin (Promega). The peptides were desalted on a StageTip C18 reverse-phase column (Thermo Scientific) and separated on an EASY-Spray column (Pepmap 3 µm, C18 15 cm × 75 µm) coupled online to a Q-Exactive Orbitrap (Thermo Scientific). The separation was performed using a flow rate of 300 nL min⁻¹ and a gradient of solvents A (0.1% FA) and B (80% ACN and 0.1% FA) as follows: 5% to 40% B for 40 min; 40% to 100% B for 5 min; 100% B for 10 min. The LC-mass-spectrometry data were processed using Proteome Discoverer software (version 1.3; Thermo Scientific) and then searched using Mascot (version 2.2.04) against the Munich Information Center for Protein Sequences barley genome database (as above). Mass spectrometry and MS/MS accuracy were 10 ppm and 20 milli mass units (mmu), respectively. In addition to oxidation of Met, variable modifications were set to include deamidation of Asn or Gln with incorporation of 18O (Asn/Gln + 0.98 D), 13C (Asn/Gln + 2.99 D), and the incorporation of one or two 18O atoms at the peptide C terminus. Carbamidomethyl Cys was set as a fixed modification. In order to be accepted as a glycopeptide, the deamidated residue containing 13C atoms had to be situated within a consensus sequence for N-glycosylation (NXS/T).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Table S1. Identification of candidate glycoproteins by Pro-Q Emerald staining in extracellular fractions of GA₃-induced aleurone layers.

Supplemental Table S2. Proteins identified in extracellular fractions of 24-h incubated aleurone layers.

Supplemental Table S3. Proteins identified in intracellular water-soluble fractions of 24-h incubated aleurone layers.

Supplemental Table S4. Identification of N-glycosylated proteins from complex samples and assignment of their glycosylation sites.

ACKNOWLEDGMENTS

We thank Birgit Andersen for excellent laboratory assistance and mass spectrometry analysis, Martin Kogel for help with microscopy experiments, and Susanne Jacobsen for valuable discussion and support.

Received November 26, 2013; accepted December 11, 2013; published December 13, 2013.

LITERATURE CITED


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