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1 Identification of Golgi-localized glycosyltransferases involved in cell wall synthesis of  
2 wheat endosperm

3

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11

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13

14 **Abstract:**

15 Plant cell walls are complex structures critical for plant fitness and valuable for human  
16 nutrition as dietary fibre and for industrial uses such as biofuel production. The cell wall  
17 polysaccharides in wheat endosperm consist of two major polymers, arabinoxylans  
18 and beta-glucans, as well as other minor components. Most of these polysaccharides  
19 are synthesized in the Golgi apparatus but the mechanisms underlying their synthesis  
20 have yet to be fully elucidated and only a few of the enzymes involved have been  
21 characterized. To identify actors involved in the wheat endosperm cell wall formation,  
22 we used a subcellular fractionation strategy to isolate Golgi-enriched fractions from  
23 endosperm harvested during active cell wall deposition. The proteins extracted from  
24 these Golgi-enriched fractions were analysed by LC-MS/MS. We report the  
25 identification of 1135 proteins among which 64 glycosyltransferases distributed in 17  
26 families. Their potential function in cell wall synthesis is discussed. In addition, we  
27 identified 63 glycosylhydrolases, some of which may be involved in cell wall  
28 remodeling. Several glycosyltransferases were validated by showing that when  
29 expressed as fusion proteins with a fluorescent reporter, they indeed accumulate in  
30 the Golgi apparatus. Our results provide new candidates potentially involved in cell  
31 wall biogenesis in wheat endosperm.

32

33 **Keywords:**

1 Glycosyltransferases / Golgi apparatus / subcellular fraction / wheat endosperm /  
2 proteomics

3 *Abbreviations:*

4 AX arabinoxylan, MLG (1-3) (1-4)  $\beta$ -D-glucan or mixed-linked glucan, GT  
5 glycosyltransferases, GH glycosylhydrolase

6

7

8 1 Introduction

9 Plant cells are surrounded by cell walls composed of polysaccharides, proteins and, in  
10 some specialized cells, various non-carbohydrate substances (e.g., lignins). Cell wall  
11 composition varies between species and between cell types and developmental stages  
12 within a given plant species. Cell wall polysaccharides have been grouped into  
13 cellulose, hemicelluloses and pectins [1]. Pectins and hemicelluloses are components  
14 of the wall 'matrix', within which the cellulose framework is embedded. The walls of  
15 most dicots and non-grass monocots (type I walls) contain about equal amounts of  
16 xyloglucan and cellulose. Grasses are noteworthy for the unusual composition of their  
17 cell walls (type II walls), which contain less pectin and xyloglucan, but more heteroxylan  
18 than the walls of other higher plants. The cell wall can accumulate extensive  
19 interconnecting networks of phenylpropanoids, particularly as the cells stop expanding  
20 and start developing secondary cell walls. The chemical structure of the components  
21 and the way that they are assembled are key factors underlying cell wall properties.  
22 These properties are important not only for plant fitness but also for the processing  
23 quality of crop materials (e.g., degradation potential for biofuel production and bread-  
24 making quality of wheat flour).

25 The cell wall composition of wheat grain endosperm is very specific, essentially  
26 composed of arabinoxylan (AX) (70%) and (1-3) (1-4)  $\beta$ -D-glucan or mixed-linked  
27 glucan (MLG) (20%) with minor amounts of cellulose (4%) and glucomannan (7%) [2,  
28 3]. The synthesis of cell wall polysaccharides is activated during wheat grain  
29 development. Drastic changes in cell wall composition and structure were observed in  
30 the developing endosperm [4-6]. In particular, a transient accumulation of (1-3)  $\beta$ -D-  
31 glucan (callose) and xyloglucan has been reported [4, 7].

32 The metabolism of cell wall polysaccharides requires the intervention of numerous  
33 enzymes. Enzymes acting on carbohydrates are listed and classified on the basis of  
34 their sequence similarities and activities in the Carbohydrate-Active enZYmes

1 database (CAZy, <http://www.cazy.org/>) [8]. Currently, CAZy contains 94  
2 glycosyltransferase (GT) families covering 122 enzymatic activities, 130  
3 glycosylhydrolase (GH) families covering 158 activities, as well as 22 families of  
4 polysaccharide lyases, 16 of carbohydrate esterases (CEs) and 64 families of  
5 carbohydrate-binding modules (CBMs). Plants are under-represented in this database,  
6 essentially because there are less fully-sequenced genomes of plant species  
7 compared to other kingdoms such as bacteria. However, in the sequenced model  
8 *Arabidopsis thaliana*, about 3.3% of all of the genes encode carbohydrate-active  
9 enzymes [9, 10].

10 Cell wall polysaccharides are synthesized from activated precursors (nucleotide  
11 sugars) by enzymatic complexes. Several groups have accumulated evidence for two  
12 major subcellular locations for cell wall polysaccharide synthesis. Cellulose microfibrils  
13 and callose are synthesized at the plasma membrane surface and the polysaccharides  
14 are deposited directly into the cell wall [11-13]. Pectin and hemicellulose are  
15 synthesized in the Golgi apparatus to then be transported to the wall [14-18].

16 The GT superfamily contains enzymes that synthesize polysaccharides such as starch  
17 and cell wall polysaccharides. GTs are also involved in the glycosylation of many other  
18 substrates such as proteins, lipids and secondary metabolites. Several GTs involved  
19 in cell wall polysaccharide synthesis have been localized in the Golgi apparatus [19-  
20 21]. GTs involved in protein glycosylation may also be localized in the Golgi apparatus  
21 and in the endoplasmic reticulum (ER) [22, 23], whereas GTs involved in starch  
22 synthesis were localized in plastids [24].

23

24 To gain access to the cellular machinery involved in the synthesis of the cell wall  
25 hemicelluloses in the wheat grain endosperm, we used a subcellular proteomic  
26 approach that targeted the Golgi apparatus to focus on the direct actors of the  
27 polysaccharide biosynthesis, i.e., the enzymes. Subcellular fractionation has been  
28 widely described across all cell types and tissues, and has been shown to be  
29 particularly efficient for organelles such as mitochondria [25] and chloroplasts [26],  
30 leading to pure fractions of these organelles. However, subcellular proteomics remains  
31 challenging for the endomembrane system. In the case of the Golgi apparatus and the  
32 ER, their similar density makes it very tedious to isolate them as pure fractions.  
33 Moreover, the large number of proteins trafficking from the ER to the Golgi apparatus  
34 inevitably leads to the identification of ER proteins, even in an ER-free Golgi

1 preparation [27]. Efficiently monitoring the subcellular fractionation and assessing the  
2 possible contamination of the isolated fractions are additional difficulties [23, 28].  
3 Finally, many GTs are membrane proteins characterized by issues such as poor  
4 solubility, low abundance and recalcitrance to classical proteomic techniques.  
5 In the present work, we have developed a subcellular strategy based on wheat  
6 endosperm Golgi fractionation from developing grain harvested at a stage where cell  
7 wall polysaccharides are accumulating. We partitioned integral membrane and water  
8 soluble proteins associated with the endomembranes. Using mass spectrometry, we  
9 identified 1135 proteins, 64 of which were predicted to be GTs involved in glycosylation  
10 processes in the wheat endosperm. We also identified 63 GHs potentially involved in  
11 the remodelling or degradation of the endosperm carbohydrates. The proteins  
12 identified comprise Golgi residents and proteins in transit in the Golgi apparatus.

13

## 14 2. Materials and methods

15

### 16 2.1. Plant materials:

17 The plants *Triticum aestivum* cv. Recital were grown in pots in a greenhouse under  
18 conditions of natural day length (UMR Amélioration des Plantes et Biotechnologies  
19 Végétales, INRA-Rennes, France). To harvest grains at defined developmental stages,  
20 individual ears were tagged at flowering. Seed development was calculated on the  
21 basis of cumulated temperature in Celsius degrees days (°D) after flowering. Grains  
22 were harvested between 250 and 275°D and endosperms were isolated manually and  
23 maintained in buffer A (250 mM sucrose, 10 mM HEPES-NaOH, pH 7.4, 1 mM EDTA,  
24 1 mM DTT and anti-protease cocktail) (Roche) on ice before further treatment.

25 For microscopy studies, the plants were grown in pots in a growth chamber at 23°C  
26 day/15°C night at 65% relative humidity under a 16h/8h day/dark photoperiod (UPR  
27 2355/IFR 87, Gif Sur Yvette, France). Wheat grains were harvested at 250°D.

28

### 29 2.2. Electron transmission microscopy

#### 30 2.2.1 Sample preparation

31 Pieces of tissue (1 mm<sup>3</sup>) were sampled from halved wheat grains. Samples were  
32 prepared as follows for morphological assay. Samples were loaded into copper hats  
33 filled with 1-hexadecene and high-pressure frozen using a HPF-EM PACT 2 freezer  
34 (Leica, <http://www.leica-microsystems.com>). The hats were immediately stored in

1 liquid nitrogen until the freeze substitution procedure was initiated. Samples were  
2 transferred to a freeze substitution automate (AFS2/FSP, Leica) and pre-cooled to -  
3 90°C. Samples were substituted in anhydrous acetone with 2% osmium tetroxyde at -  
4 90°C for 125 h. The temperature was gradually raised to -30°C for 60 h and stabilized  
5 for 12 h, then gradually raised to -4°C (34 h) and stabilized for 3 h. Samples were  
6 rinsed in anhydrous acetone several times and slowly infiltrated and embedded in  
7 Epon's resin.

8 For immunolabeling, samples were prepared as described by Chevalier et al. [29].  
9 High-pressure freezing was performed with the HPF-EM PACT I freezer. Samples  
10 were then transferred to an AFS. Samples were finally embedded in London Resin  
11 White (LRW, LRWhite R1280 HARD GRADE, London Resin Company Limited) and  
12 polymerization was performed in the AFS apparatus at -15°C under UV light for 48 h.

13

#### 14 2.2.2 Polysaccharide staining and immunolabeling

15 Periodic acid-thiosemicarbazide-silver proteinate staining (PATAg) [30] for  
16 polysaccharide detection.

17 Ultra-thin sections (80 nm) were cut from Epon's embedded samples using an  
18 ultramicrotome (MICROM MT-7000) equipped with a diamond knife and directly floated  
19 in 1% periodic acid (MERCK) solution for 30 min. They were then rinsed 5 times in  
20 deionized water before incubation in thiosemicarbazide (0.2%) (MERCK) diluted in  
21 20% acetic acid for 17 h. Sections were rinsed 5 times in decreasing acetic acid  
22 concentrations (20, 10, 5 and 2.5%), 4 times in deionized water, and stained with 1%  
23 aqueous silver proteinate (PROLABO) in the dark for 30 min. After rinsing 5 times in  
24 deionized water, sections were mounted on copper grids. Sections were examined with  
25 a JEOL 1230 transmission electron microscope with an accelerating voltage of 80 KeV  
26 (IBISA/BioGenOuest Biopolymers, Interations, Structural Biology platform (BIBS), UR  
27 1268 BIA, INRA Angers-Nantes).

#### 28 Immunolabeling

29 Ultra-thin sections (80 nm) were cut from LRW embedded samples and collected on  
30 nickel grids. Sections were incubated in a blocking solution of 3% (w/v) BSA in 20 mM  
31 PBS, pH 7.2, to block non-specific labeling for 30 min at room temperature. Sections  
32 were then incubated in a solution containing the mouse monoclonal antibody anti-AX1  
33 [31] and the rabbit polyclonal antibody raised against the pea reversibly glycosylated  
34 polypeptide 1 (anti-RGP1) (a gift from K. Dhugga), diluted 1:20 and 1:2500,

1 respectively, in 20 mM PBS supplemented with 1% BSA and 0.05% Tween-20 (Sigma-  
2 Aldrich) for 1 h at room temperature. The sections were extensively washed in buffer  
3 used for diluting the primary antibodies and then incubated 1 h at room temperature in  
4 the dark with goat-anti-mouse IgG or goat-anti rabbit conjugated with 1 nM colloidal  
5 gold complexes diluted 1:20 (v/v) in the respective buffer chosen for the primary  
6 antibodies. Labeling was then intensified with the silver enhancement kit (Aurion)  
7 according to the manufacturer's instructions. After washing, the grids were stained with  
8 2% uranyl acetate. Sections were examined with a JEOL 1230 transmission electron  
9 microscope with an accelerating voltage of 80 KeV.

### 11 2.3. Preparation of the microsomal fraction

12 Six experiments were conducted. For each experiment, approximately 20 g of  
13 endosperm were ground with a mortar and pestle on ice in buffer A and then  
14 homogenized on ice using a polytron (Kinematica AG, Dispersing and Mixing  
15 Technology) for two pulses of 7 s at 7500 rpm. The endosperm lysate was then  
16 centrifuged twice for 5 min at 2,200 g to separate cell debris. The supernatant was  
17 subsequently loaded on the top of an SW41 centrifuge tube that contained 8 mL of a  
18 18% iodixanol prepared from Optiprep solution (Sigma-Aldrich) and centrifuged at  
19 100,000 g for 2 h at 4°C (Beckman Coulter SW41 rotor). The interfacial fraction was  
20 collected as the microsome fraction.

### 22 2.4. Generation of Golgi-enriched fractions

23 Four mL of the recovered microsomal fraction were then collected from the interface  
24 and suspended in a final solution of 2.8 mM HEPES, pH 7.4, 0.28 mM EDTA, 0.28 mM  
25 DTT, 14% iodixanol and gently mixed to avoid organelle damage. These organelles  
26 were further sub-fractionated by generating an iodixanol gradient by ultracentrifuging  
27 at 200,000 g for 12 h in a Ti 70 rotor (Beckman Coulter). Fifteen fractions of 1.8 mL  
28 were harvested from the top of the gradient and their densities were determined using  
29 a refractometer (Maselli Misure; LR-01 Digital Refractometer). The protein  
30 concentration of each fraction was then determined using the NI™ Protein Assay (G-  
31 Biosciences).

### 33 2.5. Separation of organelle membranes

1 The collected fractions, microsomal and Golgi-enriched fractions, included organelles  
2 and their associated proteins. Cold deionized water was added to each fraction to  
3 reach a final volume of 22 mL in a centrifuge bottle and ultracentrifuged at 100,000 g  
4 for 2 h. The pellet consisted of organelle membranes from which membrane proteins  
5 were extracted for further identification; they will be referred to as MP (membrane  
6 proteins), MPG when issued from Golgi-enriched fractions. The supernatants of the  
7 microsomal and fraction samples were extensively dialyzed against cold water at 4°C  
8 using a semi-permeable membrane with a molecular weight cut-off of 3,500 (Spectra  
9 Pro). The dialysis water was replaced several times over the course of three days at  
10 4°C. The dialyzed proteins were then collected and lyophilized; these fractions will be  
11 referred to as AP (proteins associated with the membranes).

12

## 13 2.6. Monitoring of ER and Golgi enrichment in fractions by Western blot analysis

14 The hydrophobic nature of membrane proteins led us to choose an anionic detergent  
15 for their solubilization. Both membrane pellets and lyophilized AP were therefore  
16 solubilized in 100 µL of Tris-HCl buffer, pH 6.8, containing 2% (w/v) SDS, 10% (v/v)  
17 glycerol and bromophenol blue. Solubilized proteins were reduced by the addition of 5  
18 µL of β-mercaptoethanol and separated by SDS-PAGE on 12% acrylamide gels. After  
19 separation, gels were electroblotted onto nitrocellulose membranes (Amersham  
20 Biosciences) in 25 mM Tris, 192 mM glycine, 0.1% SDS and 20% ethanol. Semi-dry  
21 transfer was achieved at 100 V for 1.5 h.

22 Membranes were washed in PBS solution, blocked at room temperature in 5% dry milk  
23 in PBS, 0.1% Tween-20, and then washed twice for 10 min in PBS, pH 7.4, 0.1%  
24 Tween-20. Two primary antibodies were used for incubation. The polyclonal antibody  
25 anti-RGP1 (see above for immunolabeling), which recognizes a Golgi-localized  
26 protein, was used at 1:10,000 dilution for 2 h at room temperature. The antibody anti-  
27 BiP2 (Binding immunoglobulin protein2, Agrisera AB, Sweden), specific for  
28 Arabidopsis ER, was used at 1:2,000 dilution for 2 h at room temperature. After  
29 washing 2 times in PBS, 0.1% Tween-20, the secondary antibodies, consisting of  
30 peroxidase-conjugated anti-rabbit IgG, were used at 1:5,000 and 1:200,000 dilution,  
31 respectively, for 2 h at room temperature. After washing, nitrocellulose membranes  
32 were incubated in a chemiluminescent substrate (Super Signal West Dura Extended  
33 Duration Substrate, Pierce) for 6 min and dried between two paper sheets according



1 to the manufacturer's instructions. Luminescence was then detected using a camera  
2 (Luminescent Image Analyzer LAS 3000; Fuji Film).

## 3 4 2.8. Analysis of peptides by mass spectrometry

5 Proteins extracted from fractions that exhibited a high Golgi apparatus content as  
6 revealed by RGP1 were separated by SDS-PAGE on 12% acrylamide gels. The  
7 migration was performed on 2 cm of gel in order to increase the efficiency of tryptic  
8 digestion. After separation, proteins were stained with Coomassie Brilliant Blue G250  
9 (Sigma-Aldrich), as described by Devouge et al. [32].

### 10 2.8.1. Protein digestion

11 For each SDS-PAGE lane, 20 bands were manually excised from the gels to be  
12 hydrolyzed. In brief, the gel pieces were first washed three times in 50% (v/v) ACN/25  
13 mM ammonium bicarbonate, and the proteins were then reduced using DTT and  
14 alkylated with iodoacetamide prior to tryptic digestion, according to Larré et al. [33].

### 15 2.8.2. Liquid chromatography and mass spectrometry

16 Nanoscale capillary liquid chromatography-tandem mass spectrometry (LC-MS/MS)  
17 analyses of the digested proteins were performed using an Ultimate 3000 RSLC  
18 system (Dionex) coupled with an LTQ-Orbitrap VELOS mass spectrometer (Thermo  
19 Fisher). Chromatographic separation was conducted on a reverse-phase capillary  
20 column (Acclaim Pepmap C18 2 µm 100A, 75-µm i.d. x 15-cm length; Dionex) at a flow  
21 rate of 300 nL.min<sup>-1</sup>.

22 Mobile phases were composed as indicated: A (99.9% water, 0.1% formic acid), B  
23 (90% acetonitrile, 0.08% formic acid). The gradient consisted of a linear increase from  
24 4% to 45% of B in 30 min, followed by a rapid increase to 70% within 1 min.  
25 Composition was maintained at 70% B for 5 min and then decreased to 4% B for re-  
26 equilibration of the column.

27 Mass data acquisitions were performed using Xcalibur 2.1 software. Full MS scans  
28 were acquired at high resolution (FWMH 30,000) in the Orbitrap analyzer (mass-to-  
29 charge ratio (m/z): 400 to 2000), while collision-induced dissociation (CID) spectra  
30 were recorded on the five most intense ions in the linear LTQ traps.

## 31 32 2.9. Database search and interpretation

33 Raw data collected during LC-MS/MS analyses were processed into mgf format files  
34 and further searched against databanks using X!Tandem Spectrum Modeler. Protein

1 identification was achieved by confronting mass data (MS and MS/MS spectra) against  
2 four databases: (i) UniProt Knowledgebase (<http://www.uniprot.org/>) : restricted to  
3 Viridiplantae (version 2011\_03), (ii) Gene Index Databases Wheat release 12.0,  
4 August 2010 (Wheat GI) (<http://compbio.dfci.harvard.edu/tgi/>), (iii) an in-house  
5 glycosyltransferases databank (GTIDB) (<http://www.appli.nantes.inra.fr:8180/GTIDB>)  
6 [34], and (iv) a contaminant database including keratins and trypsin.

7 A database search was performed with XTandem 2008.02.01  
8 (<http://www.thegpm.org/TANDEM/>) via the X!tandem pipeline available at  
9 <http://pappso.inra.fr/bioinfo/xtandempipeline/>. Enzymatic cleavage was declared as a  
10 trypsin digestion with one possible miscleavage event. Fixed modifications of cysteine  
11 residues by iodoacetamide as well as oxidized methionins were considered. Precursor  
12 mass and fragment mass tolerance were set at 5 ppm and 0.8 Da, respectively. One  
13 missed trypsin cleavage was allowed. Identified proteins were filtered according to the  
14 following specifications: at least two peptides with an E value below 0.001 and a protein  
15 E value below  $10^{-4}$ . To take redundancy into account, proteins with at least one peptide  
16 in common were grouped.

17 Protein identifications were compared in the three databanks. In the event of  
18 identification in the three databanks, the identifier from UniProt will be preferentially  
19 reported, followed by the one from Wheat GI and, finally, the one from GTIDB. The  
20 search against the decoy database (Reverse Uniprot restricted to Viridiplantae) gave  
21 a mean false-discovery rate of 0.3% for each query when using the  $p < 0.01$   
22 significance level.

23 Tentative functional annotations were assessed by running a BLAST homology search  
24 of sequences identified from mass spectrometry against the NCBI nr database using  
25 Blast2GO (<http://www.blast2go.org>). The cut-off stringency for BLAST was  $E = 10^{-10}$ .  
26 For each protein, the best hit from each species was collected if present. The two sets  
27 of proteins (protein sequences obtained from spectrometry analysis and their best  
28 homologs) were analyzed by applying the Phobius (<http://phobius.sbc.su.se/>) and  
29 TMHMM (<http://www.cbs.dtu.dk/services/TMHMM-2.0/>) programs to predict  
30 transmembrane domains (TM) and signal peptides.

31 A refined search enabling protein identification from single peptide assignment was  
32 performed to obtain more putative GTs. This extended protein list was then blasted  
33 against the entire non-redundant sequences of the CAZy database using sequence  
34 similarity or protein domains (Pfam) association rules [35]. Those for which an ortholog

1 was found in the CAZy database (e-value < 0.01) were reported. Proteins with  
2 homology to GTs and GHs were listed, and when identified with only one peptide in  
3 MS, the quality of the peptide was checked and validated manually.  
4 For a subset of GTs and GHs, the best homolog (best Blast hit) in the rice and  
5 *Brachypodium distachyon* genomes were searched using the Blast tool of the Rice  
6 Genome Annotation website  
7 ([http://rice.plantbiology.msu.edu/analyses\\_search\\_blast.shtml](http://rice.plantbiology.msu.edu/analyses_search_blast.shtml)). Nucleotide  
8 sequences were translated with the ExPASy Translate Tool  
9 (<http://web.expasy.org/translate/>) and amino acid sequences were aligned with  
10 ClustalW2 (<http://www.ebi.ac.uk/Tools/msa/clustalw2/>).

11

## 12 2.10 Transcriptomic analysis

13 A subset of GT and GH sequences were blasted against the sequence of the wheat  
14 Affymetrix probe set using the PLEXdb Blast tool  
15 ([http://www.plexdb.org/modules/tools/plexdb\\_blast.php](http://www.plexdb.org/modules/tools/plexdb_blast.php)) [36]. The expression pattern  
16 of the corresponding “genes” was retrieved from expression studies covering wheat  
17 (whole) grain development [37] and different organs and tissues (PlaNet platform  
18 WheatNet <http://aranet.mpimp-golm.mpg.de/wheatnet>) [38]. These analyses may not  
19 discriminate all possible homologous copies within the wheat genome such as highly  
20 similar paralogs and homoeologous forms of genes from the three genomes of  
21 hexaploid wheat. Therefore, we will consider one “gene/transcript” when possibly  
22 several forms co-exist.

23

## 24 2.11. Subcellular localization

### 25 2.11.1 cDNA templates

26 Total RNA was isolated from manually dissected endosperm of wheat grains at 250°D.  
27 About 1 g of tissue was ground in liquid nitrogen and extracted in 4.5 mL of buffer (0.1  
28 M NaCl; 10 mM Tris HCl, pH 7.4, 1mM EDTA pH8; 1% SDS) with 3 mL phenol-  
29 chloroform-IAA (25:24:1). After mixing and centrifugation, the supernatant was  
30 extracted twice with 3 mL phenol-chloroform-IAA (25:24:1). The aqueous phase was  
31 precipitated by the addition of 300 µL of sodium acetate (3M, pH 5.2) and 6 mL of  
32 ethanol. RNA pellets were rinsed in 70% ethanol and dissolved in water. RNA was  
33 treated twice with the DNase set (Qiagen) and then purified through the RNeasy  
34 MinElute Cleanup kit (Qiagen), following the manufacturer’s recommendations.

1 Reverse transcription was carried out with 2 µg of total RNA from random hexamers  
2 using the Transcriptor First Strand cDNA synthesis kit (Roche).

3

#### 4 2.11.2 Constructs

5 All clones used in this study were constructed using Gateway technology (Invitrogen).  
6 The cDNA encoding the four GTs was amplified by PCR using a high fidelity KOD DNA  
7 polymerase (Novagen) with the primers listed in Supplemental Table 1. A two-step  
8 PCR procedure was carried out. A first PCR with gene-specific primers extended with  
9 part of the recombination regions attB1 and attB2 was achieved, and the resulting PCR  
10 products served as template for a second PCR, with the primers attB1 and attB2 for  
11 the GT47/Wheat GI ID: TC390543 and the GT61/UniProt ID: Q5QPY3 constructs, and  
12 the primers U5ATGSDK and U3 for the GT47/UniProt ID: E0ZPV1 and GT75/ Wheat  
13 GI ID: TC386312 constructs, to obtain the full recombination regions. Entry clones were  
14 obtained via BP reaction in pDONR-207 (Invitrogen). The binary pK7FWG2 and  
15 pK7RWG2 vectors containing the 35S promoter of cauliflower mosaic virus (CaMV)  
16 and the eGFP (enhanced Green Fluorescent Protein) or mRFP (monomeric Red  
17 Fluorescent Protein) respectively were used to construct C terminal fusions to the  
18 different GTs.

19

#### 20 2.11.3 Plant material and Agrobacterium-mediated transient expression

21 *Nicotiana tabacum* sp. plants were cultivated in a controlled growth chamber at 21°C  
22 with 14h light/10h dark for 5-6 weeks prior to Agrobacterium infiltration.

23 Transient expression of fluorescent protein fusions in tobacco leaf epidermal cells was  
24 performed as previously described by Sparkes et al. [39]. Briefly, *Agrobacterium*  
25 *tumefaciens* strains GV3101::mp90 were transformed by heat-shock with the binary  
26 plasmids. The transformants were then centrifuged and washed once with infiltration  
27 buffer (5 g/L glucose, 50 mM MES, 2 mM Na<sub>3</sub>PO<sub>4</sub>·12H<sub>2</sub>O, 100 µM acetosyringone)  
28 before being resuspended with the same buffer at an optical density (λ=600 nm) of 0.1  
29 and 0.05 for GT constructs and Golgi markers (rat sialyl transferase signal anchor  
30 sequence (ST) fused to eGFP or mRFP), respectively. Lower leaves of *Nicotiana*  
31 *tabacum* sp. plants were infiltrated with the diluted bacteria using a syringe. For co-  
32 expression, the bacteria were mixed in appropriate volumes of infiltration buffer prior  
33 to injection into the leaves. Fluorescent protein expression was studied 2-3 days post-  
34 infiltration.

1

#### 2 2.11.4 Confocal laser scanning microscopy

3 Confocal microscopy experiments were performed at the facilities of the Biopolymers-  
4 Structural Biology platform at INRA Nantes. An inverted Nikon A1 confocal laser  
5 scanning microscope was used to examine the subcellular localization of GFP and  
6 mRFP fluorescence. The samples were examined with a water-immersion x40  
7 objective. GFP and mRFP channels were acquired by simultaneous scanning using  
8 488-561 nm laser lines for excitation: emissions were collected via a photomultiplier  
9 through band-pass filters, a 500-530 nm filter and a 570-620 nm filter. Images were  
10 processed using NIS-Element Software (Nikon) and Adobe Photoshop 6.0.

11

### 12 3. Results

#### 13 3.1. Validation of the starting material and antibody used in the subcellular proteomic 14 approach

15 The wheat grain endosperm at 250°D was manually dissected and examined by  
16 fluorescence microscopy, it contains the starchy endosperm as well as the aleurone  
17 layer (Supplemental Fig. 1). At this stage, Golgi stacks are scarce compared to ER in  
18 aleurone (Fig. 1A) and in starchy endosperm (Fig. 1B) cells, making the isolation of  
19 fractions enriched in Golgi membranes challenging. Cell wall PATAg staining (Fig. 1C)  
20 and AX immunolabeling (Fig. 1D) indicate that at this stage, cell wall polysaccharides  
21 and AX, in particular, are actively produced in Golgi stacks and secreted into the wall.  
22 The antibody used to detect Golgi membranes in our fractions was initially raised  
23 against pea RGP1, a protein that has been localized in the Golgi cisternae of pea stem  
24 subapical cells [12]. We checked the reactivity of this anti-RGP1 against wheat  
25 endosperm. Immunogold labeling was detected in the Golgi apparatus of wheat  
26 aleurone cells, validating the use of anti-RGP1 to detect Golgi apparatus-enriched  
27 fractions (Fig. 2).

28

#### 29 3.2. Subcellular fractionation

30 We developed a subcellular fractionation procedure to obtain fractions enriched in  
31 Golgi apparatus from wheat endosperm. We first obtained a microsomal fraction that  
32 was further fractionated into 15 subfractions that were characterized for their relative  
33 content in Golgi stacks and ER. The fractions obtained in preliminary tests presented  
34 a low organelle content, so we decided to pool the fractions obtained from four

1 experiments. The results described in the next paragraphs were obtained from  
2 approximately 120 g of endosperm isolated from wheat grains harvested between 250  
3 and 275°D.

#### 4 5 3.2.1. Generation of the microsomal fraction containing Golgi membranes

6 In order to obtain a sufficient amount of protein for characterization and proteomic  
7 analysis from Golgi membrane-enriched fractions, six fractionations were conducted.  
8 For each fractionation experiment, we obtained approximately 20 g of endosperm from  
9 1200 wheat grains that were used to generate approximately 4 mL of microsomal  
10 fraction by concentration on an iodixanol cushion.

11 The enrichment of these fractions in organelles was examined by Western blot  
12 experiments performed with markers for two subcellular compartments, Golgi and ER  
13 with RGP1 and BiP2, respectively (Fig. 3). Western blot analysis revealed that RGP1  
14 protein was mainly present in the associated protein subfraction of the microsomal  
15 interface, whereas BiP2 protein was only found in the membrane subfraction.  
16 Accordingly, in the subsequent fractionation, the presence of BiP2 was only tested in  
17 membranes and that of RGP1 in associated protein fractions (Fig. 3).

#### 18 19 3.2.2. Generation of Golgi-enriched fractions

20 Microsomal fractions were further separated on a gradient generated from an iodixanol  
21 solution of 14%, and 15 fractions were collected along the gradient. In each  
22 experiment, the shape of the gradient was controlled by measuring the refractive index  
23 of the fractions, and a good linearity was observed from densities between 1.05 and  
24 1.1 (results not shown). The Western blots (Fig. 3) show the relative distribution of  
25 Golgi stacks and ER along the gradient fractions. The good reproducibility of these  
26 gradients led us to pool the fractions produced.

27 For protein identification, fractions revealing a high content in RGP1 and, at the same  
28 time, a low content in BiP2, were selected (Fig. 3). Fractions 6, 7 and 8 were mixed in  
29 one pool and referred to as G1, and fractions 11, 12, 13 were pooled and referred to  
30 as G2. Both G1 and G2 were further fractionated into membranes and associated  
31 proteins. The proteins extracted from the membranes pelleted from G1 and G2 will be  
32 referred to as MPG1 and MPG2, whereas the associated proteins recovered from G1  
33 and G2 will be referred to as APG1 and APG2. 1-D electrophoresis was used to reduce  
34 the sample complexity; each electrophoresis gel was cut into 20 pieces before in-gel

1 digestion and peptide extraction. These peptides were then analyzed by nano LC  
2 MS/MS.

3

### 4 3.3. Protein identification

5 The sequencing of the wheat genome is underway. To identify proteins, we therefore  
6 used UniProt restricted to Viridiplantae and, in parallel, the wheat GI database and  
7 GTIdb, an in-house database listing GT sequences from wheat, Arabidopsis and rice  
8 [34], in which each protein is assigned to a GT family according to the CAZy  
9 classification. Proteins identified from Wheat GI were annotated by running the BLAST  
10 alignment algorithm [40] against the NCBI nr database.

11

#### 12 3.3.1 Global analysis

13 First, a global analysis of all spectra generated from the four fractions was performed  
14 by combining the result files corresponding to 81 bands cut in the previously described  
15 electrophoresis experiment. This set of data generated a total of 1135 identified  
16 proteins validated with at least two peptides, and all proteins with at least one common  
17 peptide were grouped to avoid redundancy.

18 With these criteria, 668 proteins were selected (Supplemental Table 2), and the vast  
19 majority of them were identified from the Wheat GI database. Their best homologous  
20 proteins were then searched against the GenBank non-redundant protein database  
21 (NCBI), and with the exception of one sequence, homologous proteins were found with  
22 an e value  $< 10^{-10}$ . Only 20% of the sequences were identified in wheat. The others  
23 were identified by best homologs mainly found in *Hordeum vulgare* (54%), *Oryza sativa*  
24 (15%), followed by other less represented species.

25 In a second analysis the spectra were grouped on the basis of the fraction they came  
26 from, membranes (MPG) or associated (APG) Golgi proteins. Among the 668 proteins,  
27 262 were found exclusively in membrane fractions (MPG1 and MPG2), 196 in the  
28 associated protein fractions (APG1 and APG2) and 210 were found to be present in  
29 both membranes and soluble fractions. These three sub-datasets were further  
30 inspected for the presence of transmembrane domains (TM) and of signal peptides  
31 (SP) using two different software programs, Phobius and TMHMM. TM and SP  
32 predictions were examined in the sequences retrieved from the MS analysis as well as  
33 in that of their best homologs. The output of these predictions [41] is reported in Table  
34 1. Among the 262 sequences identified from the membrane fractions, 43% (112) had

1 at least one predicted TM domain. This proportion increased to 59% when the analysis  
2 was performed on their best homologs retrieved from the NCBI nr database. In the  
3 case of sequences identified exclusively in the APG fractions, the proportion of protein  
4 predicted with at least one TM drastically decreased to 4%, while among those found  
5 in both fractions, 14% were reported with at least one TM domain predicted.

### 6 7 3.3.2 Search for carbohydrate-active enzymes

8 In order to identify all putative GTs and GHs, we ran BLAST search against the entire  
9 non-redundant sequences of the CAZy database by taking, as inputs, all identified  
10 protein sequences (1135) which returned a set of 78 sequences with a pfam domain  
11 (Supplemental Table 3). Our primary targets were the GTs, but many hits were found  
12 in the other classes of carbohydrate-active enzymes. In fact, we identified 29 putative  
13 GT homologs in 11 GT families. We also identified seven sequences for carbohydrate-  
14 binding modules, five of which also included a glycosyl hydrolase "GH13" domain. We  
15 therefore decided to assign them to the GH13 family. We thus identified 46 putative  
16 GH homologs distributed in 17 GH families. In addition, one sequence belonging to  
17 Ces was retrieved. It was similar to a pectin acetylesterase from the Ces13 family.

18 In order to identify all putative GTs and GHs, we ran a new BLAST search against the  
19 CAZy database based on similarity. This secondary search resulted in identifying six  
20 additional GT families (GT2, GT30, GT41, GT61, GT77 and GT92) and two GH families  
21 (GH2 and GH12) (Supplemental Table 3). In order to avoid missing low abundant  
22 proteins we also examined the GT and GH sequences retrieved by a BLAST search  
23 on proteins identified with a single peptide. Three of them were retained as they  
24 presented a peptide with a good quality score, reported at the end of Supplemental  
25 Table 2.

26 Finally, our search resulted in an increase in the number of families identified to 17 GT  
27 families and 19 GH families (Fig. 4). Further analysis revealed that GTs and GHs were  
28 identified in a specific fraction, e.g., either GH18, GT34 in MGP (in blue, Fig. 4) or  
29 GH77, GT75 in AGP (in red, Fig. 4). Surprisingly, some of them were found in both  
30 fractions.

31 Among the GT and GH families identified, the GT2, GT34, GT47, GT48, GT61, GT75,  
32 GH3, GH9 and GH17 were reported in the literature to contain members potentially  
33 involved in cell wall metabolism. The wheat sequences identified in these families were



1 assigned to their best BLAST hit in the genome of rice and *Brachypodium distachyon*  
2 and their putative or characterized functions are reported in Table 2.

### 3 4 3.4 Expression pattern of a subset of candidate genes

5 We analyzed the expression pattern of the subset listed in Table 2 by mining published  
6 and publicly available transcriptomic data. In Supplemental Fig. 2, the results show  
7 that several “genes” such as members of the GT34, GT61, GH3, GH9 and GH17  
8 families exhibit a pattern of expression relatively specific to the grain. This result  
9 confirmed the importance of choosing the target tissue of interest as starting material  
10 for our analysis.

### 11 12 3.5 Experimental validation of the Golgi localization for three chosen GTs

13 To validate the Golgi localization of candidates identified with the subcellular proteomic  
14 approach, we chose GT families known for their potential involvement in cell wall  
15 biosynthesis. In particular, we selected GTs for which a complete nucleotide coding  
16 sequence was available at the time we did our analysis, and validated them *in silico*  
17 after nucleotide and amino acid sequence alignments with their best BLAST hit in rice  
18 and *Brachypodium distachyon*. The coding sequences corresponding to the  
19 GT75/Wheat GI ID: TC386312, GT47/UniProt ID: E0ZPV1, GT61/UniProt ID: Q5QPY3  
20 (GenBank ID AK333763.1), and GT47/Wheat GI ID: TC390543 were inspected and  
21 the first three were found to be complete. In the case of GT47/Wheat GI ID: TC390543,  
22 sequence alignments with the best *Brachypodium* homolog and with wheat ESTs  
23 revealed a missing nucleotide in the coding sequence that shifted the reading frame  
24 (Supplemental Document 1). The *in silico* validated coding sequences of the three GTs  
25 were cloned using cDNA obtained from the endosperm of wheat grain at 250°D. The  
26 intracellular localization of the GT proteins was determined by transiently expressing  
27 them as fluorescent protein fusions in tobacco leaf epidermal cells. GT75/Wheat GI ID:  
28 TC386312, GT47/UniProt ID: E0ZPV1, GT61/UniProt ID: Q5QPY3, and GT47/Wheat  
29 GI ID: TC390543 were all expressed as small and mobile bodies in the cells, highly  
30 similar to the Golgi bodies. Co-expression of the GTs with a Golgi marker confirmed  
31 their localization in the Golgi apparatus (Fig. 5).

## 32 33 4. Discussion

1 The wheat endosperm cell wall composition is unusual and relatively simple, and  
2 appears to be a good model to reach enzymes implicated in the biosynthesis of their  
3 constituting polymers. It is thought that part of these enzymes and especially the GTs  
4 are located in the Golgi apparatus. Many of these GTs are membrane proteins present  
5 in low abundance; they were not detected in global proteomic analyses of the wheat  
6 grain endosperm [42-44]. The challenge of this work was to identify carbohydrate-  
7 active wheat enzymes at the proteomic level with a special focus on GTs potentially  
8 involved in the synthesis of cell wall polysaccharides in the grain endosperm. Additional  
9 difficulties had to be overcome. Only a few Golgi apparatus are visible in the starchy  
10 endosperm at 250°D. We chose this developmental stage because the endosperm is  
11 then easily isolated and because there is an active deposition of cell wall  
12 polysaccharides (in particular, the AX starts to accumulate), whereas grain filling with  
13 storage compounds (starch and storage proteins) is only beginning. Another difficulty  
14 lays in the fact that wheat proteins and especially wheat GTs are not well represented  
15 in protein databases.

16 We developed a procedure for the isolation of Golgi-enriched fractions adapted from  
17 that of Graham [45] and Sadowski et al. [46] followed by 1D protein analysis, in-gel  
18 digestion and mass spectrometry analysis of the resulting peptides.

19

#### 20 Isolation of Golgi-enriched fractions

21 The microsomal fraction we obtained revealed an enrichment in Golgi stacks and ER.  
22 The BiP2 was used as an ER marker and was only revealed in the membranes of our  
23 microsomal fraction, corroborating the results obtained by Mitra et al. [47] who  
24 identified BiP2 in the membrane sub-proteome of whole Arabidopsis seedlings.

25 RGP1 was used as a Golgi marker. We, however, detected RGP in both the soluble  
26 and the membrane subfractions of the microsome, which confirmed the presence of  
27 Golgi stacks in the microsomal fraction and corroborates previous results that report  
28 that RGPs have been found in both the membrane and soluble fractions of many  
29 species [48, 49]. A RGP was identified in the associated fraction (Wheat GI ID:  
30 TC386312) in our proteomic analysis.

31 Variation of the relative enrichment in Golgi or ER was observed along the gradient,  
32 allowing us to collect the most Golgi-enriched fractions for protein identification. The  
33 presence of BiP2 in these fractions can be either due to ER contamination or to a  
34 possible localization in Golgi as reported recently in Arabidopsis cell culture [50].

## 1 Proteomic analysis

2 The subcellular fractionation and LC–MS/MS analysis of wheat endosperm resulted in  
3 the identification of 1135 unique proteins (at least two peptides with an E value < 0.001)  
4 that, after grouping, gave 668 non-redundant proteins, 262 from the membrane  
5 fractions and 196 from the associated fractions, meaning that 210 were found in both  
6 fractions.

### 7 Membrane and associated proteins

8 Proteins only detected in the membrane fractions are expected to be strongly linked to  
9 the membranes (integral membrane proteins and proteins anchored to the membrane),  
10 whereas proteins within the membrane-associated fractions may be proteins in transit  
11 within the endomembrane system and proteins linked to membrane proteins.

12 The proteins identified in both fractions may be due to a strong association between a  
13 protein and a membrane protein within a complex where some but not all proteins are  
14 integral membrane proteins as already described by Zeng et al. [28].

15 Furthermore, as recently reported by Atmodjo et al. [51], the transmembrane domain  
16 of some type II proteins are cleaved in the Golgi apparatus, although the processed  
17 proteins are retained in the Golgi apparatus within protein complexes. We cannot  
18 exclude the possibility that for some of the proteins, the dual location may reflect a  
19 contamination due to insufficient membrane washing, although the fractions were  
20 extensively diluted (30 times) before membrane pelleting.

21 The prediction analysis of the transmembrane domain reflects this distribution: only  
22 several percent of the proteins within the associated fraction are predicted to contain  
23 a TM domain, whereas half of the proteins found only in the membrane fractions are  
24 predicted to contain such a domain. This percentage would undoubtedly increase if the  
25 wheat protein sequences were complete since many Golgi proteins have their  
26 transmembrane domain in the N-terminal region.

### 27 Golgi localization

28 Many proteins transit via the endomembrane system (ER and Golgi) before being  
29 directed to their final destination. In particular, secreted proteins undergo glycosylation  
30 in the ER and the Golgi apparatus, and it has been postulated that about 50% of plant  
31 proteins could be glycosylated [52]. Some proteins are retained in the Golgi apparatus  
32 where they carry on their function.

33 In our Golgi-enriched fractions, we identified proteins that are not supposed to be Golgi  
34 residents: abundant proteins (histones, ribosomal proteins, ribulose-1,5-bis-phosphate

1 carboxylase/oxygenase) that contaminate our fractions, as well as GTs such as the  
2 callose synthase (Wheat GI ID: TC451164), most probably localized in the plasma  
3 membrane and the starch synthase (UniProt ID: Q43654|SSY1\_WHEAT) in plastids.  
4 In addition to their transit in the Golgi apparatus, proteins can be recycled via the Golgi  
5 apparatus. For example, the cellulose synthase and possibly the callose synthase,  
6 both located in the plasma membrane, could be stored in the Golgi apparatus or  
7 recycled to be directed where needed [50, 53]. In order to validate the Golgi location  
8 of some of the GTs identified, four of them were co-expressed with a Golgi marker in  
9 tobacco cells.

10

## 11 Identification of proteins of interest

12

### 13 Storage proteins

14 Our study focused on one specific developmental stage that corresponds to the very  
15 early stage of storage deposition (starch and protein). The quality of our samples was  
16 confirmed by the low amount of storage proteins found by our analysis. Less than 1%  
17 of the identified proteins were storage proteins - five gliadins and three glutenins -  
18 which are secretory proteins synthesized and folded into the ER before being  
19 deposited into organelles known as protein bodies [54]. However, these proteins were  
20 also detected in Golgi-derived vesicles early in the grain development, indicating that  
21 the Golgi apparatus is also involved in prolamin transport [55, 56].

### 22 Carbohydrate-active enzymes

23 Within the numerous proteins identified, it was predicted that 78 sequences would be  
24 active on carbohydrates, among them, proteins from 17 GT families (Fig. 4).

25 Several of the GTs identified are clearly not involved in cell wall synthesis. GT4 and  
26 GT5 are enzymes that catalyze the metabolism of sucrose (sucrose synthase UniProt  
27 ID: Q8W1W4) and starch (starch synthase UniProt ID: Q43654). Interestingly, they  
28 were only identified in the associated fraction. Members of the GT66 family (STT3  
29 homolog GenBank ID: AAT69659) were implicated in protein N-glycosylation and  
30 localized in the ER [23].

31

### 32 GTs potentially involved in cell wall polysaccharide biosynthesis

33 Other GTs identified in our proteomic analysis have already been implicated in cell wall  
34 synthesis. This is the case for the GT2 CslF6 protein of wheat (Wheat GI ID:

1 TC390697). Homologs were identified as MLG glucan synthases following the insertion  
2 of rice *CsIF* genes into the *Arabidopsis thaliana* genome, which resulted in the  
3 detection of grass-specific MLG in the walls of the transgenic lines [57]. The wheat  
4 gene *CsIF6* was identified and silenced by RNA interference, resulting in a decrease  
5 of the MLG content in wheat grain endosperm [58].

6 Conversely, we identified GTs known to be involved in AX synthesis. The GT47-13  
7 protein UniProt ID: E0ZPV1\_WHEAT was found in a complex with  
8 glucurono(arabino)xylan synthase activity in a wheat seedling microsomal fraction [28].  
9 Although no transmembrane domain is predicted for this protein, we localized it in the  
10 Golgi apparatus. The TaGT47-13 protein is the potential ortholog of Arabidopsis IRX10  
11 and IRX10-L [59, 60], which have been implicated in the elongation of the xylan  
12 backbone.

13 RGP proteins are members of the GT75 family, although their role in glycosylation  
14 remains to be proven. Indeed, several recombinant rice and *Arabidopsis thaliana*  
15 RGPs were found to have a UDP-Arap mutase (UAM) activity, which is required to  
16 convert the UDP-arabinopyranose into UDP-arabinofuranose [61, 62]. Down-  
17 regulation of rice UAM reduces the proportion of arabinofuranose in rice cell walls [63].  
18 Zeng et al. [28] detected a GT75 protein (UniProt ID: Q9ZR33) in the  
19 glucurono(arabino)xylan complex that they isolated from a wheat seedling microsomal  
20 fraction. We identified three GT75 in the wheat endosperm (Wheat GI ID: TC386312,  
21 Wheat GI ID: TC371253, Wheat GI ID: TC399081), the sequence of Wheat GI ID:  
22 TC386312 is highly similar but not identical to UniProt ID: Q9ZR33. Our fusion  
23 experiment revealed that Wheat GI ID: TC386312 was localized in the Golgi apparatus,  
24 despite a lack of transmembrane domain.

25 Very recently, several rice and wheat GT61 of the subgroup A were shown to be xylan  
26 O3 arabinosyltransferases (XATs) [19]. We identified three different GT61 of the  
27 subgroup A in the wheat endosperm (Table 2) and localized one of them, the UniProt  
28 ID: Q5QPY3 corresponding to XAT1, in the Golgi apparatus. Transcriptomic analysis  
29 showed that it seems specifically expressed in the endosperm. A GT61 from the  
30 subgroup B (UniProt ID: Q5QPZ5\_WHEAT) was also found in our analysis (Table 2).  
31 The corresponding gene exhibits an expression pattern specific to the grain  
32 endosperm and floral bracts before anthesis (Supplemental Fig. 2). This GT61 may be  
33 an arabinosyltransferase required for the double substitution of xylose residues in the

1 endosperm AX at position O2 and O3. It is noteworthy that no GT61 was identified in  
2 the glucurono(arabino)xylan complex isolated from wheat seedlings by Zeng et al. [28].  
3 A GT48 protein was revealed in our analysis. The GT48 family is known to include the  
4 glucan synthase-like proteins (GSL) or callose synthase (CalS) [64, 65]. Callose plays  
5 an important role in plant development and in response to stress. Callose is deposited  
6 at cell plates during cytokinesis. It accumulates transiently in the wheat endosperm  
7 early in the development [4]. Voigt et al. [66] identified eight *GSL* genes in wheat and  
8 studied their expression in different organs. The protein we detected in the endosperm  
9 at this stage (Wheat GI ID: TC451164) corresponds to the *TaGSL12* gene.

10 It was very recently revealed that members of the *CsID* family have mannan synthase  
11 activity [67]. Mannans are present in the wheat endosperm. Nemeth et al. [58]  
12 produced RNA interference lines targeting the wheat *CsID2* gene potentially encoding  
13 the *CsID2* protein that we detected by proteomic analysis (Wheat GI ID: TC394160).  
14 They only investigated a potential effect on AX and MLG. Thus, it is possible that the  
15 wheat *CsID2* has a mannan synthase activity. Virtually nothing is known about the  
16 structure, i.e., the possible substituents of mannans in wheat grain. Among the proteins  
17 we identified in our proteomic analysis, several GT34 share similarity with a  
18 galactomannan galactosyltransferase identified in lotus seeds [68]. The transcripts  
19 encoding the GT34 Wheat GI ID: TC370787 were detected in the grain but not in the  
20 vegetative organs investigated, and the transcript level increases steeply in the  
21 developing grain (Supplemental Figure 2). Mannans have a storage function in the  
22 seeds of lotus and many other species [69]. Nothing is known about their function in  
23 the wheat grain. Another activity was described for members of the GT34 family.  
24 Several *Arabidopsis* genes of this family were found to encode xyloglucan  
25 xylosyltransferase [70-72]. Xyloglucan were recently detected transiently in the wheat  
26 endosperm [7].

27 Another interesting GT identified is the GT47 Wheat GI ID: TC390543 (best hit in rice  
28 LOC\_Os11g03410), a protein containing a single transmembrane domain and that we  
29 localized in the Golgi apparatus. The GT47 family is encoded by 39 genes in  
30 *Arabidopsis* [73]. Phylogenetic analyses of the GT47 family revealed that  
31 LOC\_Os11g03410 (and Wheat GI ID: TC390543) is in a different subgroup of GT47  
32 (subgroup C) than GT47-13 (UniProt ID: E0ZPV1) and *Arabidopsis* IRX10-10L, which  
33 are in subgroup A according to the Zhong and Ye classification [7, 73]. Subgroup C  
34 contains several rice and *Arabidopsis* proteins. Only two of these proteins (ARAD1 and

1 ARAD2) have an ascribed function based on mutant characterization: they are putative  
2 arabinosyltransferases acting on pectin [74]. Wheat GI ID: TC390543 and its rice  
3 ortholog LOC\_Os11g03410 are not really close to the ARAD proteins, and although  
4 pectin is a component of the rice endosperm [75], no pectin has been reported in the  
5 wheat grain endosperm [3]. Thus, it is possible that the enzyme encoded by Wheat GI  
6 ID: TC390543 has a different activity.

7 We achieved our aim by identifying GTs potentially involved in wheat endosperm cell  
8 wall synthesis through a proteomic approach.

9

10 GHs potentially involved in cell wall polysaccharide remodelling

11 Surprisingly, we also identified many GH proteins that we did not expect to be located  
12 within the Golgi apparatus. To our knowledge, Golgi location was described for some  
13 GHs, such as members of the families GH38 (Wheat GI ID: TC416977) and GH47  
14 (Wheat GI ID: TC393888), which would be an alpha-mannosidase involved in protein  
15 glycosylation [23, 50, 76]. It is thought that most of the other GHs are secreted. Many  
16 GHs are glycosylated and probably pass through the Golgi apparatus. Half of these  
17 GHs were identified in the associated fraction and only a few of them were predicted  
18 to contain a TM domain, consolidating the idea that we identified them during their  
19 transit through the endomembrane system.

20 The GH proteins identified in the endosperm fell into 18 of the 34 families that are  
21 represented in the rice, Arabidopsis and Brachypodium genomes [77].

22 The putative function of some of the individual GHs was searched by studying the  
23 annotation of their best homologs in other species. However, the occurrence within the  
24 same family of numerous enzymatic activities makes the task of identifying the  
25 potential activity of individual GHs difficult. Some GHs identified here are clearly not  
26 involved in cell wall polysaccharide modification/turnover. In fact, the GH14 proteins  
27 are beta-amylases (UniProt ID: Q7X9M2\_WHEAT). Other families include enzymatic  
28 activities that could modify cell walls.

29 The GH17 family contains many characterized activities, including  $\beta$ -1,3-glucanases  
30 (EC 3.2.1.39) that specifically degrade callose, as shown in Arabidopsis  
31 plasmodesmata [78]. We identified several GH17 in our experiment. BJ242908 is  
32 similar to the rice putative  $\beta$ -1,3-glucanase (UniProt ID: Q9ZNZ1) and may be related  
33 to the early and transient deposition of callose in the grain cell wall that is degraded  
34 during development since callose is not detected in the mature grain endosperm. The

1 expression of the gene encoding GenBank ID: BJ242908 seems to be specific to the  
2 grain endosperm and increases during grain development (Supplemental Fig. 2).  
3 Fused with GFP, the callase identified in the plasmodesmata was localized in the  
4 membrane and the ER [78].

5 Among the GH9, some were predicted to have an endo- $\beta$ -glucanase activity. We  
6 identified several GH9 sequences (GenBank ID: CD888519\_WHEAT, Wheat GI ID:  
7 TC382188, Wheat GI ID: TC442952). Best hit results by similarity search are endo- $\beta$ -  
8 glucanase (cellulase) implicated in leaf and fruit abscission in *Prunus persica* ppEG1,  
9 *Citrus sinensis* and *Malus x domestica* [79]. In *Arabidopsis thaliana*, KORRIGAN is a  
10 GH9 endo- $\beta$ -glucanase that was proposed to play a role in cellulose synthesis [80-82].  
11 The cell wall of wheat grain endosperm contains a minor amount of cellulose [2]. The  
12 transcript corresponding to Wheat GI ID: TC382188 seems to be specific to the grain  
13 endosperm and its expression increases during grain development (Supplemental Fig.  
14 2).

15 Several GH3 were revealed in our analysis. Several activities were reported for  
16 enzymes within the GH3 family. Among the GH3 identified in our screen, Wheat GI ID:  
17 TC369308 is homolog to the barley  $\beta$ -D-glucan exohydrolase isoenzyme Exoll  
18 (UniProt ID: Q42835\_HORVU), and Wheat GI ID: TC422719 and Wheat GI ID:  
19 TC384225 are homologs to a xylan  $\beta$ -D-xylosidase characterized in barley (Genbank  
20 ID: AAK38482). Transcripts encoding the putative xylosidases are well represented in  
21 the endosperm, and the abundance of Wheat GI ID: TC422719 increases considerably  
22 during early development (Supplemental Fig. 2). Genbank ID: CK196562 is homolog  
23 to the arabinofuranosidase/xylosidase identified in germinating barley (Lee et al. 2003).  
24 The substrates of all these enzymes are most likely to be MLGs and AXs, the two main  
25 components of the wheat endosperm cell wall. At the onset of their synthesis, the AXs  
26 of wheat endosperm are highly substituted with arabinose and later in the endosperm  
27 development, the level of AX substitution decreases [4, 5, 83, 84]. Toole et al. [84]  
28 proposed that a GH with arabinofuranohydrolase or arabinofuranosidase activity would  
29 remove arabinose residues to decrease the level of AX substitution. Recent studies  
30 using films of pure AXs as models of endosperm cell walls revealed the influence of  
31 AX structure on water diffusion [85, 86]. Higher water diffusion was obtained for films  
32 made with highly-substituted AXs, compared to films made with lowly-substituted AXs.  
33 This suggests that AX-decreased substitution may have a physiological function during



1 development, probably in the regulation of water/solute diffusion during grain  
2 filling/dessication.

3

4 In summary, using this subcellular proteomic approach, we have successfully identified  
5 over a thousand proteins from wheat starchy endosperm with many new enzymes  
6 probably involved in cell wall biogenesis. They are members of families already  
7 assumed to be involved in cell wall synthesis and modification or degradation, but  
8 information about their function is still lacking. The expression pattern of the candidates  
9 reveals that all of them were expressed in the wheat endosperm, with a few of them  
10 exhibiting an expression profile specific to the endosperm. Synthesis of cell wall  
11 hemicellulose takes place in the Golgi apparatus. We report here GTs potentially  
12 involved in this mechanism, four of which were localized in the Golgi apparatus.

13

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15

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25

26

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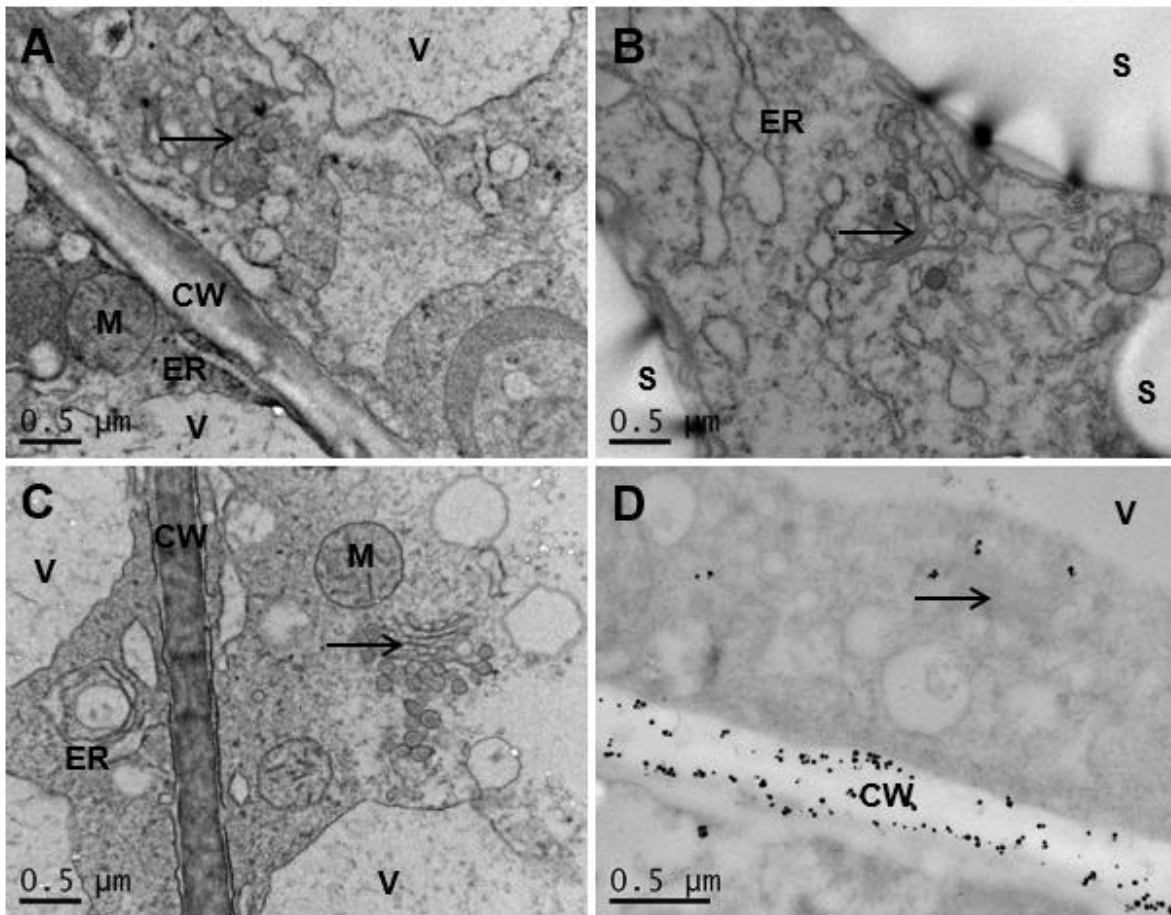
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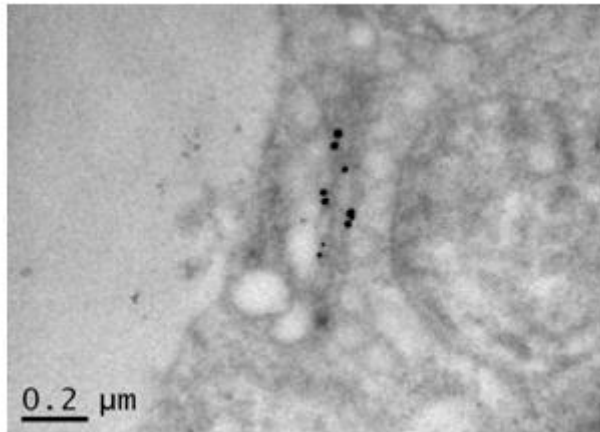
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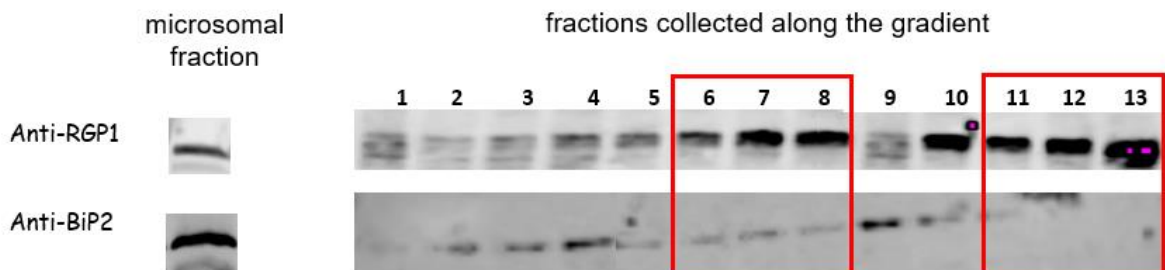
1  
 2 Figure 1: Transmission electron micrographs of developing wheat grain at 250°C. A  
 3 view of the cytoplasm showing different organelles in aleurone cells (A) and in the  
 4 starchy endosperm (B). The aleurone cells exhibit dense granular cytoplasm with  
 5 numerous vacuoles, ER and mitochondria. A few Golgi stacks are observed. In the  
 6 starchy endosperm, ER tubules, some of which show dilation, are abundant, whereas  
 7 Golgi structures with stacked cisternae are barely visible. PATAg staining of aleurone  
 8 cell walls (C). PATAg-stained materials are detected within the cell walls and in Golgi-  
 9 derived vesicles. Immunogold labeling of arabinoxylan in aleurone cells (D).  
 10 Arabinoxylan epitopes are visualized in the cell wall. Labeling is mainly localized in the  
 11 inner region of the primary cell walls and found over Golgi stacks.  
 12 CW, cell wall; ER, endoplasmic reticulum; M, mitochondria; S, starch granule, V,  
 13 vacuole; arrow, Golgi stacks.

14



1  
2 Figure 2: Immunogold localization of RGP1 in aleurone cells. Gold particles occur over  
3 Golgi stacks but none over other cell structures.

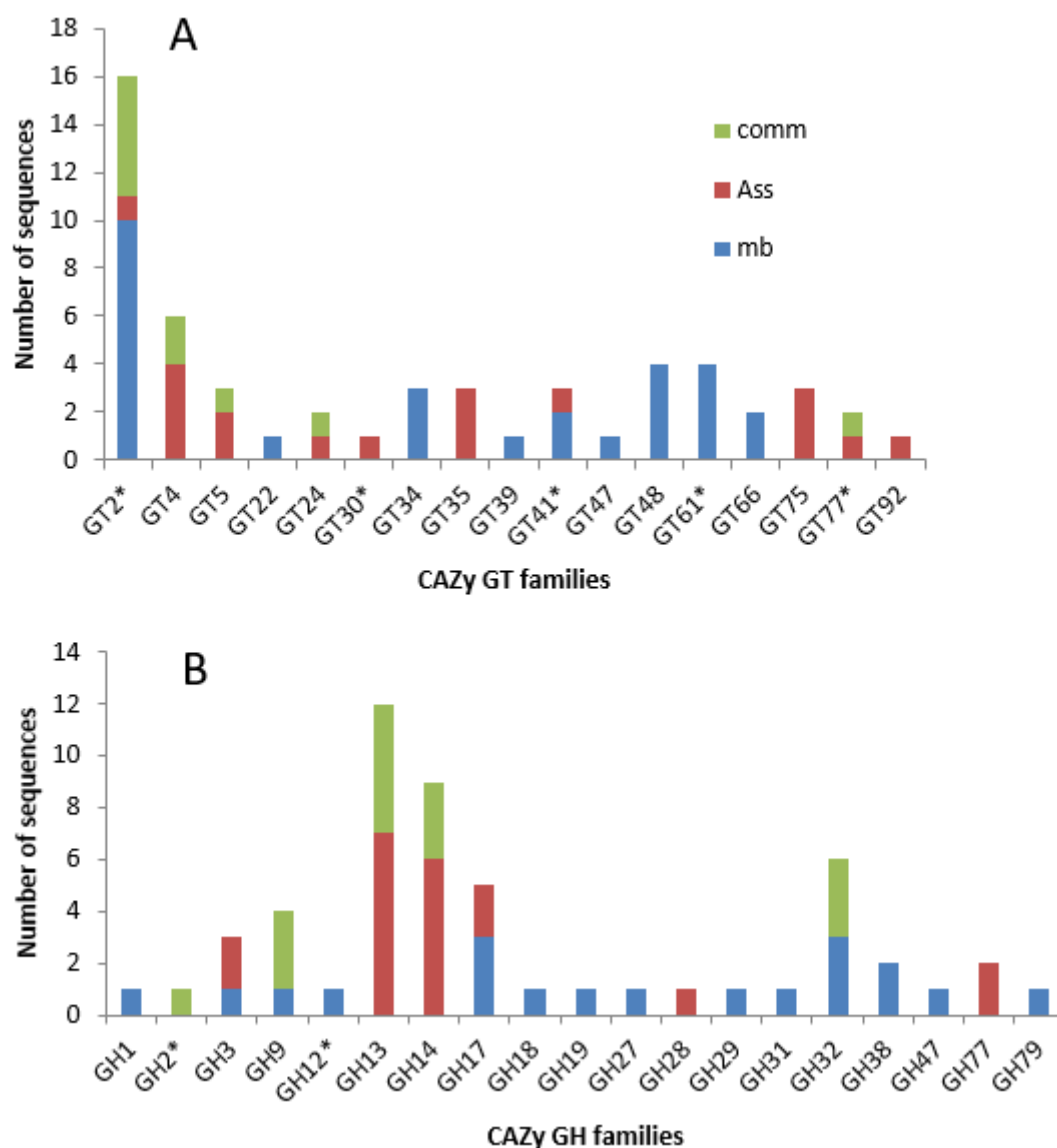
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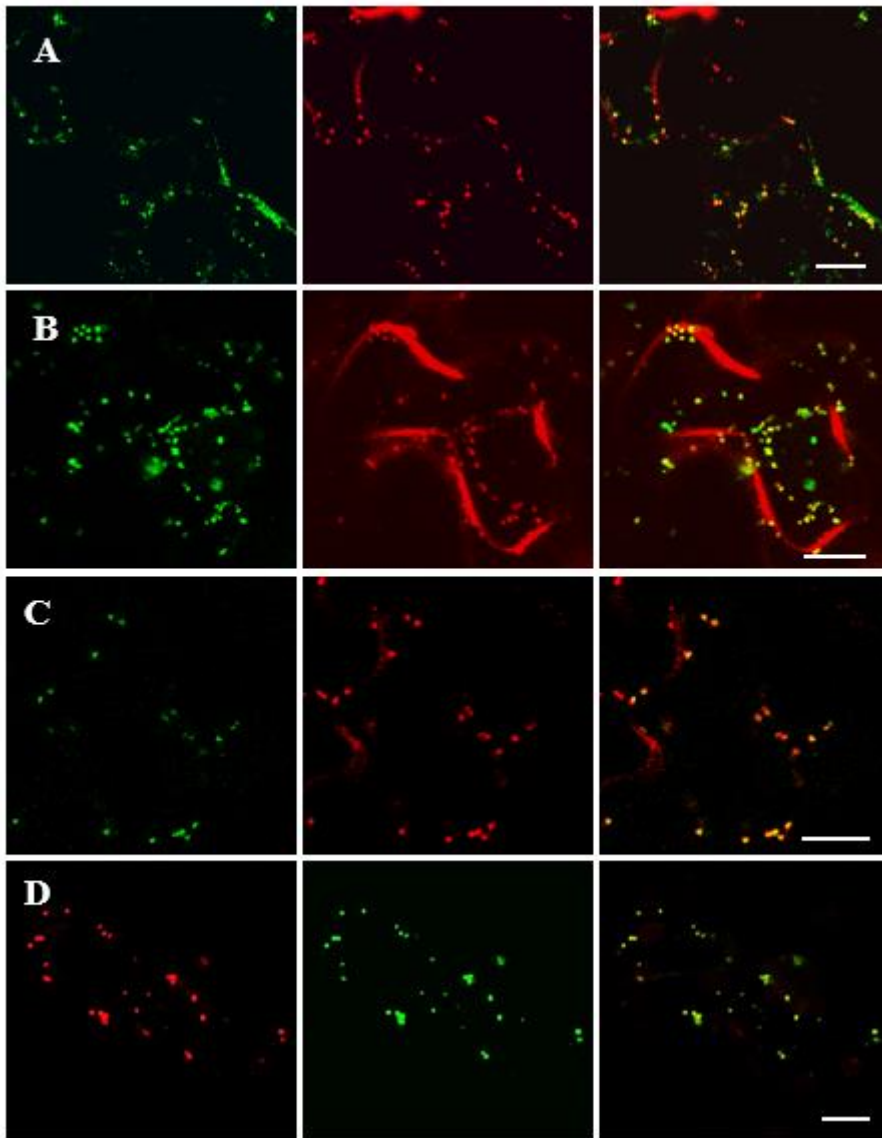
5  
6 Figure 3: Presence of Golgi apparatus and ER in microsomal fraction and fractions  
7 collected along the gradient revealed by western blot with antibodies against Golgi  
8 protein (Anti-RGP1 antibody) and ER protein (Anti-BiP2 antibody) as indicated. The  
9 anti-RGP1 results presented here were obtained on the associated protein fractions  
10 and those of Anti-BiP2 on membrane protein fractions.

11





1  
 2 Figure 4: Distribution of blasted sequences in (A) glycosyltransferase (GT) families and  
 3 in (B) glycosylhydrolase (GH) families; number of hits obtained with the Pfam domain  
 4 except for GTs and GHs labeled with a star for which the hits were obtained only by  
 5 sequence similarity. Colors correspond to the fractions in which the sequences were  
 6 identified: blue: present in membrane fractions; red: present in associated fractions;  
 7 green: present in both fractions.  
 8



1  
 2 Figure 5: The subcellular localization of some glycosyltransferases fused to fluorescent  
 3 proteins in tobacco epidermal cells. Confocal images showing eGFP (green) and  
 4 mRFP (red) fusion proteins coexpressed in tobacco leaf epidermal cells 2-3 days after  
 5 agrobacterium infiltration. Images of cells expressing GT47-TC390543-eGFP (green)  
 6 co-expressed with a Golgi marker, ST-mRFP (red) and merged image (A), GT75-  
 7 TC386312-eGFP (green) co-expressed with ST-mRFP (red) and merged image (B),  
 8 GT61-Q5QPY3-eGFP (green) co-expressed with ST-mRFP (red) and merged image  
 9 (C), GT47-E0ZPV1-mRFP (red) co-expressed with ST-eGFP (green) and merged  
 10 image (D), Scale bars = 10  $\mu$ m.

11

			Phobius			TMHMM	
		Total number of sequences	TM	SP	TM (%)	TM	TM (%)
Membrane (MPG)	sequences	262	106	39	40.6	112	42.9
	best Homologues	262	137	87	52.5	152	58.2
Associated (APG)	sequences	196	7	20	3.6	12	6.2
	best Homologues	196	5	34	2.6	12	6.2
both fractions	sequences	210	30	36	14.3	35	16.7
	best Homologues	210	35	63	16.7	35	16.7

**Table 1** : Prediction of signal peptides (SP) and transmembrane domains (TM) using Phobius and TMHMM.

1

	proteomic hit	group	Families	best blast hit rice	E-value	annotation/putative fonction (rice)	brachypodium hit	hit activity in family and/or subgroup
Wheat GI ID:	TC390697	359	GT2	LOC_Os08g06380	0	CSLPB - cellulose synthase-like family F; beta1,3;1,4 glucan synthase	Bradi1g25130	MLG synthase
Wheat GI ID:	TC394180	242	GT2	LOC_Os08g02180	0	CSLD2 - cellulose synthase-like family D	Bradi1g50170	mannan synthase
Wheat GI ID:	TC370787	535	GT34	LOC_Os02g49140	1.0e-77	glycosyltransferase, putative	Bradi3g58510	galactomannan galactosylTase
Wheat GI ID:	TC413183	958	GT34	LOC_Os11g34390	1.4e-75	glycosyltransferase, putative	Bradi4g16870	galactomannan galactosylTase, xyloG xylosylTase
Wheat GI ID:	TC390543	444	GT47	LOC_Os11g03410	2.0e-209	exostosin family domain containing protein	Bradi4g26150	subgoup C arabinosyltransferase acting on pectins
UniProt ID:	E0ZPV1	1091	GT47	LOC_Os01g70190	4.3e-219	exostosin family domain containing protein	Bradi2g59400	subgroup A xylan xylosylTase (xylan synthase?)
Wheat GI ID:	TC451164	137	GT48	LOC_Os06g02280	6.2e-125	callose synthase, putative	Bradi1g51780	callose synthase
UniProt ID:	Q6QPY3	299	GT81	LOC_Os02g22850.2	2.5e-184	glycosyltransferase, putative	Bradi3g11300	XAT1 clade A xylan O3 arabinosyltransferase
Wheat GI ID:	TC437977	597	GT81	LOC_Os02g22850	1.5e-142	glycosyltransferase, putative	Bradi3g11300	clade A xylan O3 arabinosyltransferase
UniProt ID:	DBL9S8	863	GT81	LOC_Os01g02930	3.5e-130	glycosyltransferase, putative	Bradi2g01420	clade A xylan O3 arabinosyltransferase
UniProt ID:	Q6QPZ5	511	GT81	LOC_Os01g72810	6.4e-170	glycosyltransferase, putative	Bradi2g61230	clade B unknow n function
Wheat GI ID:	TC386312	203.1	GT75	LOC_Os03g40270	1.1e-198	alpha-1,4-glucan-protein synthase, putative	Bradi1g15050	RGRUDP-Aramputase
Wheat GI ID:	TC371253	203.2	GT75	LOC_Os03g40270	5.1e-133	alpha-1,4-glucan-protein synthase, putative	Bradi1g15050	RGRUDP-Aramputase
Wheat GI ID:	TC399081	203.3	GT75	LOC_Os07g41380	1.3e-178	alpha-1,4-glucan-protein synthase, putative	Bradi1g21990	RGRUDP-Aramputase
Wheat GI ID:	TC384225	289	GH3	LOC_Os02g51620	2.3e-235	glycosyl hydrolase family3 protein, putative	Bradi3g59020	xylan B-D-xylosidase
Wheat GI ID:	TC422719	594	GH3	LOC_Os11g18730	1.2e-179	glycosyl hydrolase family3 protein, putative	Bradi4g20200	xylan B-D-xylosidase
Wheat GI ID:	TC389308	197	GH3	LOC_Os03g53800	1.2e-240	beta glucosidase	Bradi1g08580	MLG endohydrolase
GenBank ID:	OK198562	1195	GH3	LOC_Os04g54810	0	beta-D-xylosidase, putative	Bradi6g23470	arabino furanosidase
GenBank ID:	CD888519	655	GH9	LOC_Os05g03840	1.8e-28	endoglucanase, putative	Bradi2g38380	endo-1,4 beta-glucanase activity (cellulase)
Wheat GI ID:	TC382188	84	GH9	LOC_Os04g36610	1.4e-83	endoglucanase, putative	Bradi6g11310	endo-1,4 beta-glucanase activity
Wheat GI ID:	TC442952	317	GH9	LOC_Os04g36610	2.1e-45	endoglucanase, putative	Bradi6g11310	endo-1,4 beta-glucanase activity
GenBank ID:	BJ242908	641	GH17	LOC_Os01g71810	5.9e-69	glycosyl hydrolases family 17, putative	Bradi2g60490	beta 1,3 glucanase calase
Wheat GI ID:	TC395632	237	GH17	LOC_Os01g71380	7.7e-86	glycosyl hydrolases family 17, putative	Bradi2g60490	beta 1,3 glucanase calase
Wheat GI ID:	TC394973	468	GH17	LOC_Os01g71680	4.9e-37	glycosyl hydrolases family 17, putative	Bradi2g60490	beta 1,3 glucanase calase

**Table 2** : List of candidate GTs and GHs for a role in cell wall metabolism in the developing wheat endosperm

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