

Central Functions of the Lumenal and Peripheral Thylakoid Proteome of Arabidopsis Determined by Experimentation and Genome-Wide Prediction

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Experimental proteome analysis was combined with a genome-wide prediction screen to characterize the protein content of the thylakoid lumen of Arabidopsis chloroplasts. Soluble thylakoid proteins were separated by two-dimensional electrophoresis and identified by mass spectrometry. The identities of 81 proteins were established, and N termini were sequenced to validate localization prediction. Gene annotation of the identified proteins was corrected by experimental data, and an interesting case of alternative splicing was discovered. Expression of a surprising number of paralogs was detected. Expression of five isomerases of different classes suggests strong (un)folding activity in the thylakoid lumen. These isomerases possibly are connected to a network of peripheral and lumenal proteins involved in antioxidative response, including peroxiredoxins, m-type thioredoxins, and a lumenal ascorbate peroxidase. Characteristics of the experimentally identified lumenal proteins and their orthologs were used for a genome-wide prediction of the lumenal proteome. Lumenal proteins with a typical twin-arginine translocation motif were predicted with good accuracy and sensitivity and included additional isomerases and proteases. Thus, prime functions of the lumenal proteome include assistance in the folding and proteolysis of thylakoid proteins as well as protection against oxidative stress. Many of the predicted lumenal proteins must be present at concentrations at least 10,000-fold lower than proteins of the photosynthetic apparatus.

INTRODUCTION

Recently, the genome of the dicotyledon Arabidopsis was sequenced completely, and 25,498 genes were annotated (Arabidopsis Genome Initiative, 2000). The availability of the Arabidopsis genome now allows the classification of proteins according to predicted domain(s) and subcellular localization. Importantly, the sequenced genome also makes it possible to take advantage of the dramatic improvements in biological mass spectrometry to study gene expression directly at the protein level and to determine protein localization and post-translational modifications efficiently in a systematic manner. The improved mass accuracy, mass

resolution, and sensitivity of the latest generation of mass spectrometers allow the rapid identification of picomole to femtomole amounts of proteins and peptides. Examples of excellent reviews on mass spectrometry and their application in biology include those by Jensen et al. (1998), Pandey and Mann (2000), and Yates (2000) and references in the study by Blackstock and Mann (2000). Combined with rapidly expanding plant genome information, these technical improvements are expected to have a profound effect on plant biology (for discussion, see van Wijk, 2001).

A significant subset of the Arabidopsis proteome is localized in different organelles (e.g., the mitochondria and different types of plastids). Plastids are predicted to contain between 10 to 15% of the nucleus-encoded gene products, corresponding to ~2500 to 3500 proteins, indicating the importance of this organelle in the plant cell (Arabidopsis Genome Initiative, 2000). This prediction is based on the presence of an N-terminal chloroplast transit peptide (cTP) in the nucleus-encoded chloroplast proteins (except for

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proteins localized in the chloroplast outer membrane) using the cellular localization program TargetP (Emanuelsson et al., 2000). After synthesis in the cytosol, the cTP is recognized at the chloroplast envelope and the precursor protein is translocated into the chloroplast, followed by processing of the cTP. Proteins then are directed into the inner envelope membrane, into the thylakoid membranes, or remain in the chloroplast stroma. The inner envelope proteins, the peripheral thylakoid proteins located at the stromal site of the thylakoid membrane, and most of the integral thylakoid membrane proteins have no additional N-terminal transit peptides. In contrast, proteins located in the thylakoid lumen are targeted and translocated via a second transit peptide, the luminal transit peptide (ITP), which is located directly C-terminal of the cTP. A few small thylakoid proteins with a single transmembrane (TM) domain also have an ITP (Thompson et al., 1998, 1999). Luminal proteins are involved in a number of well-characterized functions, such as water splitting, electron transport, and the violaxanthin cycle. However, the function of most newly discovered luminal proteins is not clear (Kieselbach et al., 1998; Peltier et al., 2000).

The ITPs show strong similarities to bacterial signal peptides and can be divided into a charged N-terminal domain, a hydrophobic core, and a more polar C-terminal domain that ends with short chain residues at the -3 and -1 positions relative to the terminal cleavage site (von Heijne et al., 1989). Soluble luminal proteins with ITPs are translocated by at least two different mechanisms involving two different sets of proteins, whereas the single TM proteins with ITPs appear to insert spontaneously (reviewed by Dalbey and Robinson, 1999; Keegstra and Cline, 1999). Proteins translocated via the so-called Sec pathway require ATP and cpSecA/Y (Knott and Robinson, 1994; Settles et al., 1997; Mori et al., 1999) and probably cpSecE (Schuenemann et al., 1999; Froderberg et al., 2001). Proteins translocated via the Δ pH or twin-arginine translocation (TAT) pathway require neither soluble factors nor GTP or ATP, but they do require a transthylakoid proton gradient and several TAT proteins. The TAT substrates also require a specific TAT motif, RRx-h-h, where R is arginine and h is a hydrophobic residue (Walker et al., 1999; Robinson and Bolhuis, 2001; for an exception, see Summer et al., 2000).

In a recent study, we began to systematically identify proteins in thylakoids from pea using two-dimensional (2-D) electrophoresis and mass spectrometry (MS) (Peltier et al., 2000; van Wijk, 2000). Because little genomic or protein sequence information is available for pea, most of the proteins were identified based on homology with proteins in Arabidopsis. The nucleus-encoded proteins identified on the 2-D gels were used to test the prediction for chloroplast localization and the transit peptide by the software programs ChloroP (Emanuelsson et al., 1999) and SignalP (Nielsen et al., 1997, 1999). The program SignalP (<http://www.cbs.dtu.dk/services/SignalP/>) was developed originally for the prediction of cleavable signal peptides of secretory proteins in

bacteria and eukaryotes, but it also predicted ITPs as well as their cleavage sites with good success (Peltier et al., 2000).

With the complete annotated genome of Arabidopsis now available, we set out to identify a larger fraction of luminal proteins and to extend the set of experimentally determined N termini. We expected that this would provide parameters and thresholds for a genome-wide theoretical prediction of the luminal proteome using modified versions of the cellular localization programs TargetP (an update of ChloroP) and SignalP as a basis. We also thoroughly verified the gene annotation of all experimentally identified luminal proteins and demonstrate generic strategies for reannotation. The molar ratio between the most abundant and least abundant identified proteins was 5×10^4 , and many of the predicted luminal proteins are expected to have expression levels that are 10^5 - to 10^6 -fold below those of the most abundant luminal proteins.

RESULTS

Experimental Identification of Luminal and Weakly Bound Peripheral Proteins by 2-D Electrophoresis and MS

To purify the luminal proteome, thylakoids from Arabidopsis were purified carefully using differential centrifugation, linear Percoll gradients, and multiple washing steps. Subsequently, the purified thylakoids were disrupted by Yeda press treatment. The luminal proteome then was separated by 2-D electrophoresis using immobilized pH gradient (IPG) strips in the first dimension (pI 4 to 7 and 6 to 11) and high-resolution Tricine gradient gels in the second dimension. Both analytical (250 μ g of protein) and preparative (1.5 to 2 mg of protein) 2-D gels were generated and were stained with silver or Coomassie blue, respectively. Figures 1A and 1B show the analytical 2-D gels of the purified lumen at the two pI ranges. For identification of the proteins, the strategy used was similar to that described by Peltier et al. (2000). In summary, protein spots were picked from the 2-D gels, washed, digested with the site-specific protease trypsin, and extracted. The masses of the extracted peptides were determined by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) MS and/or sequenced by nano-electrospray ionization tandem mass spectrometry (nano-ESI/MS/MS) and followed by identification using different search engines. N-terminal sequences of ~ 30 identified proteins were determined by Edman degradation to validate ITP prediction.

The proteins identified on the 2-D gels and the summarizing data of the identification process are listed in the tables, grouped according to the predicted or most likely cellular localization. For each protein identified, we have listed the percentage of sequence coverage obtained by MALDI-TOF MS (at 50 ppm mass accuracy), one of the several internal sequences determined by tandem MS, and the N-terminal sequence as determined by Edman sequencing. In addition,

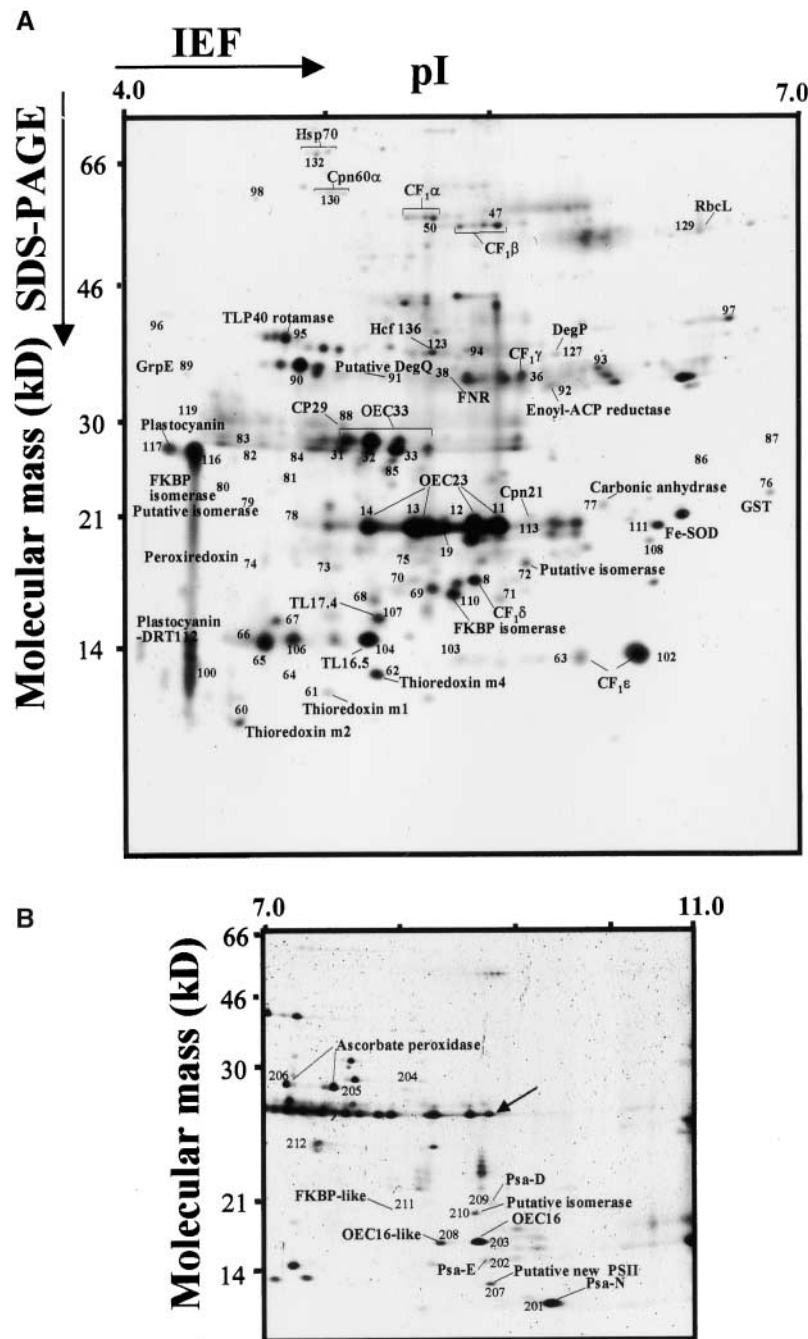


Figure 1. Silver-Stained 2-D Electrophoresis Maps of Luminal Proteins of Arabidopsis (ecotype Columbia).

Fractions strongly enriched for thylakoid luminal proteins isolated from Arabidopsis were separated by 2-D electrophoresis with denaturing iso-electric focusing (IEF) on immobilized pH gradients in the first dimension between pH 4 and 7 (**A**) or between pH 7 and 11 (**B**) and Tricine-PAGE in the second dimension (**A**) and (**B**). Gels were calibrated for molecular mass (in kD) and pI (in pH units) by internal (pH and mass) and external (mass) standards. The protein spot numbers refer to the spot numbers listed in Tables 1 to 4. For a selected number of spots, the identity (in addition to the number) also has been listed on the 2-D maps. The map in (**B**) also includes the internal standards for the pI, as indicated by the arrow. To more easily compare the Arabidopsis maps with those constructed for pea (as described by Peltier et al., 2000), spot numbers are identical for orthologous pairs.

ACP, acyl carrier protein; FNR, ferredoxin-NADP reductase; GST, glutathione S-transferase; PSII, photosystem II; SOD, superoxide dismutase.

Table 1. Identification of 30 Luminal Arabidopsis Proteins from the 2-D Electrophoresis Gels Shown in Figure 1

Spot No.	Molecular Mass (kD)	pI	Identity ^a	Accession No. ^b	MALDI Percent Coverage at 50 ppm ^c	MS/MS Sequence ^d	Targeting Pathway	Localization and Cleavage Site Prediction ^e						
								TargetP	Predotar	Euk	SignalP		N Terminus	
											Gram ⁻	Gram ⁺		
203	16.7	9.3	OEC16	4583542 (At4g21280)	44		TAT	C2/44	P 1	75-VLA-DA	74-AVL-AD	77-ADA-IS	77-ISIKV ^f	
208	16.7	9.1	OEC16-like	7267278 (At4g05180)	37		TAT	C3/48	P 0.7		84-AEA-IP	84-AEA-IP	84-IPIKV ^f	
11, 12, 13, 14	20.5	5.7 to 6.0	OEC23	7443217 (At1g06680)	40		TAT	C2/31	P 0.6	77-ADA-AY	77-ADA-AY	77-ADA-AY	77-AYGEA ^{g,h}	
201	12.1	9.6	Psa-N	1709825 (At5g64040)	35		TAT	C5/81	P 0.9	84-ASA-NA	86-ANA-GV	84-ASA-NA	84-NAGV ^f	
123	36.7	5.6	Hcf136	6016183 (At5g23120)	47		TAT	C2/60	P 0.9	78-ARA-DE	78-ARA-DE	78-ARA-DE	78-DEQLS ^g	
205 to 206	30.4	7.2 to 7.6	Putative ascorbate peroxidase	7267543 (At4g09010)	32		TAT	C3/36	P 1			82-AKA-AD	82-ADLIQ ^g	
207	13.2	9.3	Putative new photosystem II protein	2809245 (At1g03600)	33		TAT	C2/67	P 1	68-VSA-AE	68-VSA-AE	68-VSA-AE	68-AEED ^g	
91	33.6	5.4	Putative DegQ protease (DegP5)	2832642 (At4g18370)	11		TAT	C3/26	P 0.5		10-AFS-SI	71-AIA-LE	71-XEQX ^h	
19	19.4	5.6	Putative protein OEC23 related	2244908 (At4g15510)	23		TAT	C1/75	M 0.7	104-AFA-ST	104-AFA-ST	54-QSA-KS	104-STPV ^f	
71	17.2	6.0	Not annotated OEC23 related	AC007171		SLDQFGSPQFVAD (3)	TAT (KR)	C4/34	P 0.8	71-AKS-ME	26-SHH-KI	69-NKA-KS		
204	31.4	8.5	Putative protein OEC23 related	7019666 (At3g55330)	31		TAT	C1/26	Neither	74-SFA-AE	74-SFA-AE	74-SFA-AE	74-AESKK ^g	
212	26.3	7.6	Putative protein OEC23 related	7485407 (At2g39470)	54		TAT	C3/72	Neither	73-LLA-EE	73-LLA-EE	73-LLA-EE		
108	19.0	6.5	Putative protein OEC23 related	2829916 (At1g77090)	33		TAT	C4/35	M 1		63-ALA-FP	63-ALA-FP	70-VVKKQ ^h	
111	19.5	6.5	Putative protein OEC23 related	7594543 (At3g56650)		GSTVWLFVVSATEK (7)	TAT	C1/75	P 0.9		66-ISA-AR	66-ISA-AR	67-REVEV ^{g,h}	
210	17.3	9.3	Putative isomerase	6143884 (At3g10060)	50		TAT	C2/56	Both	73-ASG-IL	82-AEA-VS	82-AEA-VS		
110	17.3	5.7	Putative FKBP isomerase	2289010 (At2g43560)	50		TAT	C4/48	Neither	59-AAG-LP	57-AYA-AG	57-AYA-AG	AGLPP ^f	
80	25.4	4.7	Similar to FKBP isomerase	7543908 (At5g13410)	24		TAT	C1/29	P 0.6		31-VAA-RL	17-ALA-GT	89-SQFAD ^f	
104	16.0	5.2	Putative protein	2262151 (At4g02530)	23		TAT	C1/38	P 0.9		58-LVA-IG	58-LVA-IG	73-AILEA ^g	
70	17.8	5.5	Putative protein OEC23 related	6642664 (At1g76450)		VEAFAETLVSGDLR (3)	TAT	C2/32	P 0.8	80-AFA-ET	80-AFA-ET	80-AFA-ET	80-ETNAS ^f	
117	28.9	4.2	Plastocyanin 2	130261 (At1g76100)	23		Sec + other	C1/32	P 0.9		72-AMA-ME	72-AMA-ME	72-AMA-ME	72-MEVLL ^f
90, 100, 116	14 to 35	4.4 to 4.9	Plastocyanin 1 DRT112	1169201 (At1g20340)	29	IEVLLGGGDSGLAFIPNDFSIAK (4)	Sec + other	C1/75	P 1		68-AMA-IE	68-AMA-IE	68-IEVLL ^h	
31 to 33	28.5	5.1 to 5.6	OEC33	3286693 (At5g66570)	55		Sec + other	C2/26	P 0.8	85-ASA-EG	85-ASA-EG	85-ASA-EG	85-EGAPK ^f	
			OEC33-like	4835233 (At3g50820)	37		Sec + other	C2/28	P 0.9	84-AGA-EG	84-AGA-EG	84-AGA-EG	84-EGAPK ^f	
127	36.5	6.2	DegP1	2565436 (At3g27925)	31		Sec + other	C3/35	P 1	100-VES-AS	99-AVE-SE	100-VES-AS	103-FVVS ^f	
95	38.5	4.8	TLP40 rotamase	6016707 (At3g01480)	53		Sec + other	C1/75	P 1	82-AHA-VA	82-AHA-VA	82-AHA-VA	92-VLISG ^f	
211	21.2	8.5	FKBP-like protein	6686798 (At4g39710)	32		Sec + other	C3/74	M 0.6	73-ADA-TR	73-ADA-TR	73-ADA-TR		
107	16.7	5.3	Putative protein pentapeptide repeat	6226234 (At5g53490)	36	AFVGNIGQADGVYDKPLDLR (1)	Sec + other	C4/65	P 0.7	77-VIA-AN	77-VIA-AN	77-VIA-AN	77-ANQRL ^g	
103	15.0	5.7	Putative protein pentapeptide repeat	2344892 (At2g44920)	31		Sec + other	C1/56	P 0.8	81-ALA-FK	81-ALA-FK	81-ALA-FK	81-FKGGG ^g	
106	16.0	4.8	Putative protein	8809586 (At5g52970)		VLAQNPVTPGLAIK (4)	Sec + other	C4/83	P 1	75-ADA-KV	75-ADA-KV	75-ADA-KV	75-KVGVN ^g	
73	18.2	5.0	Putative protein	3776572 (At1g54780)	44		Sec + other	C2/53	P 1	84-ALA-SE	84-ALA-SE	84-ALA-SE	84-SEFN ^h	
68	17.1	5.3	Putative protein	4455236 (At4g24930)		FWLEDTPYGR (2)	Sec + other	C4/66	M 0.5	63-ALA-IP	63-ALA-IP	63-ALA-IP	63-IPSLS ^f	

Proteins were identified by MALDI-TOF MS and/or nanoESI/MS/MS. N termini of all proteins were determined by Edman sequencing. All accession numbers in italics showed mis-assignments in the database and have been corrected.

^aIdentity given in NCBI completed by domain prediction found by Pfam, Blocks, Prints, Prodom, or Phi-Blast.

^bAccession numbers in NCBI and in MIPS (in parentheses). All accession numbers in italics showed misassignments in the database and have been corrected.

^cPercentage of coverage at 50 ppm for the MALDI-TOF peptides.

^dSequence tag identified by ESI/MS/MS; additional sequence tags were obtained (numbers listed in parentheses) but are not shown.

^eLocalization, reliability class, and cleavage site prediction by TargetP (C for chloroplast), localization and score by Predotar (P for plastid, M for mitochondria), cleavage site prediction of the luminal transit peptide by three versions of SignalP developed for eukaryotes (Euk), Gram-negative or Gram-positive bacteria, and in boldface when this prediction fits with the N-terminal Edman sequence.

^fN-terminal Edman sequence tag found in the literature or in public databases.

^gN-terminal Edman sequence tag identified in our previous study of pea.

^hN-terminal Edman sequence tag identified in this study.

the localization predictions by TargetP and Predotar, as well as the predictions of ITPs obtained using SignalP in three different search modes (eukaryotic, Gram-positive, and Gram-negative bacteria), are indicated. A prediction of the targeting pathway (TAT or Sec + other) through the thylakoid membrane based on the characteristics of the ITP also is listed. This extensive prediction of location and cleavage sites was performed to set thresholds and parameters for the theoretical genome-wide prediction of the luminal proteome, as presented in the second half of this article. It is important to stress that we did not attempt to experimentally identify all luminal proteins (an impossible task); rather, we sought to assemble a set of luminal proteins that would allow determination of the parameters and thresholds for theoretical prediction. This has the advantage that an overview of the total potential luminal proteome and function can be obtained, including proteins of very low abundance or proteins that are expressed only under specific conditions or in different types of plastids (see Discussion).

Protein Identities, Function, and Expression of Isoforms and Paralogs

Eighty-one proteins on the 2-D gels were identified with high confidence. Thirty of those are very clearly luminal proteins,

based on characteristic presequences and on very good matches between the experimentally determined N termini and the predicted ITPs (Table 1). Nineteen of those luminal proteins have signal sequences that are typical for TAT substrates. An additional 12 proteins, listed in Table 2, have fairly clear features of ITPs, and for several of them the experimentally determined N terminus corresponded to the predicted luminal cleavage site. However, because of an ambiguous Edman sequence tag, because the ITP cleavage site is unusual (see further below), or because of contrasting localization information in the literature, they were not added to Table 1 and their true localization needs to be determined by additional experimentation.

To quantify the expression levels of the proteins in both pea (Figure 2 from Peltier et al., 2000) and Arabidopsis, silver-stained and Coomassie blue-stained gels (pl 4 to 7 maps) were analyzed using the 2-D software package Melanie. All expression levels were calculated on a molar basis and normalized within each gel to the expression of Hcf136, because Hcf136 spots could be well quantified on both the Coomassie blue- and silver-stained maps (Figure 2). Spots are shown only if they were detected in duplicate on both the Coomassie blue- and silver-stained gels (thus, four gels for each spot). The expression levels of the most abundant proteins (the oxygen-evolving complex [OEC] proteins and plastocyanin [PC]) were ~10,000-fold higher than those of

Table 2. Identification of 12 Arabidopsis Proteins from the 2-D Electrophoresis Gels Shown in Figure 1

								Localization and Cleavage Site Prediction ^a					
Spot No.	Molecular Mass (kD)	pI	Identity ^a	Accession No. ^b	MALDI Percent Coverage at 50 ppm ^c	MS/MS Sequence ^d	Targeting Pathway	SignalP					
								TargetP	Predotar	Euk	Gram ⁻	Gram ⁺	N Terminus
74	18.4	4.7	2-Cys peroxiredoxin	7242491 (At3g11630)		APDFEAEAVFDQEFIK (1)	TAT	C2/48	P 1		69-AQA-DD	69-AQA-DD	
68	17.1	5.3	Peroxioredoxin-like	7529720 (At3g52960)		YAILADDGVVK (3)	TAT	C1/56	P 0.9		71-VTI-SI	73-ASI-SV	73-SVXXK ^f
79	22.2	4.7	Putative isomerase	7287985 (At3g60370)	38		TAT	C5/42	Neither	53-SSS-AK	54-SSA-KT	54-SSA-KT	
89	34.6	4.3	Chaperone GrpE	4583546 (At5g11710)	45		TAT	C2/64	Neither	67-ASG-EA	68-SGE-AE	67-ASG-EA	
69	17.6	5.6	Putative protein	5840783 (At5g58250)		VHFLVANAK (7)	Sec + other	C1/52	M 0.8	18-AAA-CR	58-KTA-AT	58-KTA-AT	
60	12.1	4.6	Thioredoxin m1	2809238 (At1g03680)	35		Sec + other	C2/48	Neither			69-CEA-QD	
61	13.3	5.0	Thioredoxin m2	4206206 (At4g03520)	34		Sec + other	C1/56	Neither			75-CEA-QE	73-EAQET ^f
62	14.3	5.3	Thioredoxin m4	6539614 (At3g15360)	21		Sec + other	C1/82	P 0.9	27-SSA-AP	27-SSA-AP	69-RIA-RG	76-EAQDT ^f
77	21.3	6.3	Carbonic anhydrase	14343 (At3g01500)	46		Sec + other	C2/32	P 0.8	19-SQS-SL	64-VFA-AP	64-VFA-AP	66-APXIA ^f
131	75.9	4.9	Hsp70	7441883 (At4g24280)	21		Sec + other	C1/92	P 0.8			93-AVA-AM	
130	61.8	5.0	Cpn60	2506276 (At2g28000)	24		Sec + other	C2/45	P 1		14-VLC-SS	49-ANV-KE	
113	20.6	6.2	Cpn21	4127456 (At5g20720)	23		Sec + other	C2/50	P 1		19-SLA-SL	92-AQS-KP	

The locations of these proteins are either in the thylakoid lumen or peripherally attached to the stroma side of the thylakoid membrane. All proteins were identified by MALDI-TOF MS and/or nano-ESI/MS/MS. N termini for four proteins were determined by Edman sequencing.

^aIdentity given in NCBI completed by domain prediction found by Pfam, Blocks, Prints, Prodom, or Phi-Blast.

^bAccession numbers in NCBI and in MIPS (in parentheses).

^cPercentage of coverage at 50 ppm for the MALDI-TOF peptides.

^dSequence tag identified by ESI/MS/MS; additional sequences were identified (numbers in parentheses) but are not shown.

^eLocalization, reliability class, and cleavage site prediction by TargetP (C for chloroplast), localization and score by Predotar (P for plastid, M for mitochondria), cleavage site prediction of the luminal transit peptide by three versions of SignalP developed for eukaryotes (Euk), Gram-negative or Gram-positive bacteria, and in boldface when this prediction fits with the N-terminal Edman sequence.

^fN-terminal Edman sequence tag identified in this study.

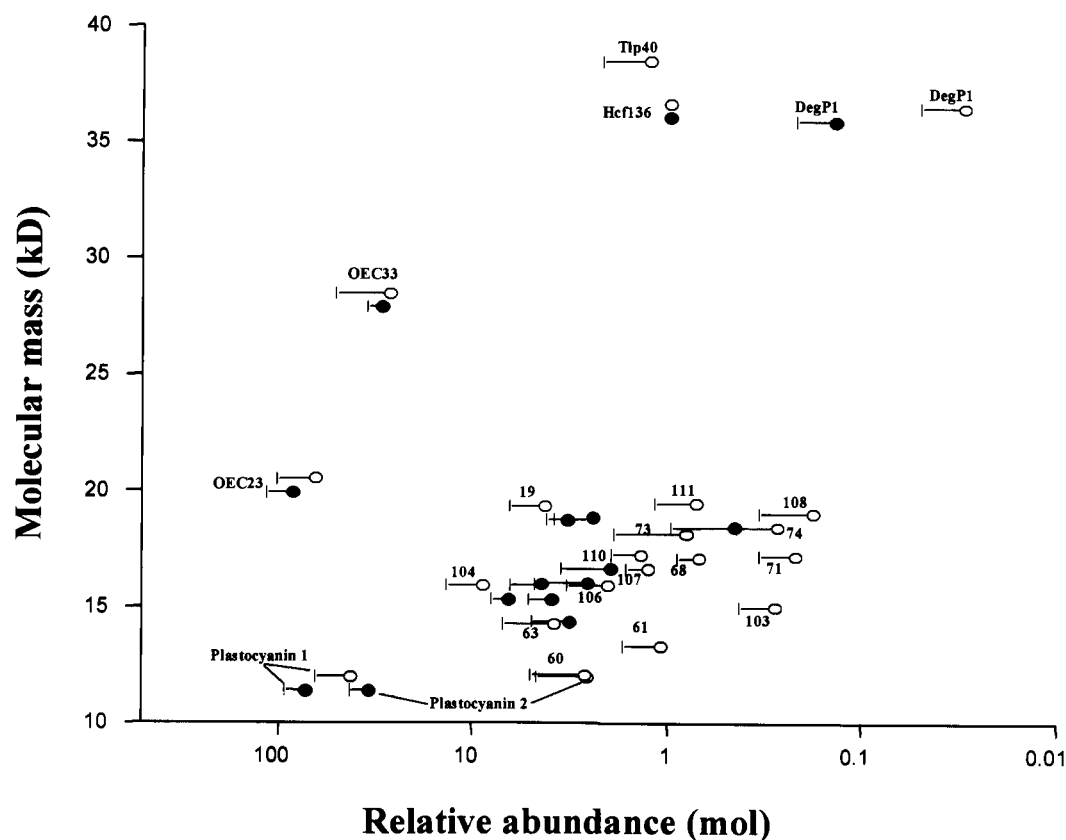


Figure 2. Relative Expression Levels of Thylakoid Proteins in Pea and Arabidopsis.

Quantification of expression levels of luminal and other thylakoid proteins from Arabidopsis proteins (closed symbols) and pea proteins (open symbols) calculated on a molar basis and normalized to the expression level of Hcf136. Duplicate Coomassie blue-stained and duplo silver-stained 2-D gels of two independent experiments with a pH range of 4 to 7 were analyzed. Standard errors ($n = 4$) are indicated, and the x axis is in log scale.

the least abundant proteins (see below for details). In general, the expression levels for pea and Arabidopsis corresponded very well (Figure 2).

Apart from OEC23, seven weakly related OEC23 paralogs were identified on the 2-D gels (Table 1). In our earlier study of pea thylakoids, we also observed expression of an eighth paralog (spot 22). Sequence tags for some of the OEC23 paralogs (spots 19, 108, 111, and 204) also were identified on our previous maps of the pea lumen; however, at the time, we were not always able to find the corresponding genes or the OEC23 domains. The expression levels of the paralogs were 20- to 300-fold lower than that of OEC23. Thus, OEC23 is by far the most strongly expressed protein in this family. Database searching identified a ninth OEC-like protein that was 88% identical to OEC23 (accession number At2g30790). However, expression data could not be found from expressed sequence tags (ESTs) or at the protein level. Sequence analysis of this OEC23-related gene

showed a frameshift (as a result of a base pair loss) at the end of the first exon in the ITP, possibly preventing protein expression.

For three other luminal proteins (OEC16, OEC33, and PC) and several peripheral proteins (fibrillins, RNA binding protein 29, and m-type thioredoxins), one or more very closely related paralogs were identified (Figures 3A and 3B). The difference in expression levels within each group of paralogs was on the order of a factor 5 to 25. Alignment of pairs of these closely related paralogs showed that the presequences were less conserved than were the mature proteins, with an average 63% identity for the presequences and 78% for the mature proteins (Figures 3A and 3B). It is likely that these paralogs originate from gene duplications (see Discussion).

Interestingly, a multigene family of five luminal isomerases (Table 1) and one potential luminal isomerases (Table 2) was identified experimentally, of which only TLP40

was identified previously (Fulgosi et al., 1998). The relationship for this isomerase family is shown in Figure 4 in the form of a rooted tree. Two of the isomerases (spots 210 and 211) are basic proteins (pI 8.3 and 9.3) and were expressed at much lower levels than was OEC16 (Figure 1B). The expression levels of TLP40 and spot 110 were quite similar to the expression of Hcf136 (Figure 2), whereas spot 79 was expressed at very low levels (<0.01).

Other luminal or putative luminal proteins include several known chaperones (GrpE, HSP70, Cpn60, and Cpn21), the assembly factor Hcf136 (Meurer et al., 1998), the protease DegP1 described by Itzhaki et al. (1998), and carbonic anhydrase. In addition, eight luminal proteins (Table 1) and two more potential luminal proteins (Table 2) with unknown functions were identified. We identified three m-type thioredoxins (m1, m2, and m4) as well as two 2-Cys peroxiredoxins and sequenced the N termini of m2 and m4 and one of the peroxiredoxins. Recently, these five proteins were implicated directly in antioxidant defense (Baier and Dietz, 1999; Baier et al., 2000; Issakidis-Bourguet et al., 2001), and their localization is not clear but probably is peripheral on the stromal side of the thylakoid membrane (Baier et al., 2000).

In addition to the luminal proteins, 32 proteins were identified that are either (1) peripheral thylakoid proteins attached to the stromal side and are part of photosynthetic complexes (two Psa proteins and five CF₁ proteins) or (2) associated with the thylakoid membranes as part of their function (six ribosomal proteins, four RNA binding proteins, a ribosome recycling factor, six fibrillins, Fe-superoxide dismutase, and ClpS1) and very low levels of the two ribulose-1,5-bisphosphate carboxylase/oxygenase subunits, which are the most abundant stromal proteins in the chloroplast (Table 3). Interestingly, we also identified ROC4, a cyclophilin with peptidyl-prolyl *cis-trans* isomerase activity, which was determined earlier to be in chloroplasts by protein gel blotting (Lippuner et al., 1994). The copurification with the luminal proteins indicates that they interact with the thyla-

koid membrane but that they are released easily by either the Yeda press or the sonication used to open the luminal compartment. Because we did not observe any significant amount of the very abundant Calvin cycle enzymes, it is clear that this set of 32 proteins interacts with the membrane as part of their function.

Finally, low levels of six nonchloroplast proteins from different cellular locations were identified (Table 4). All six proteins are known to be very abundant in each of these locations. Several of these proteins were identified exclusively by sequence tags and not by MALDI-TOF MS. Together, these six proteins contribute <0.5% of total mass, indicative of the low level of contamination from nonchloroplast locations.

Correction of Gene Annotations by MS Analysis and Comparison with ESTs

We systematically analyzed the proteins listed in Table 1 and several others for quality of gene annotation in the Arabidopsis database. Significant errors were detected in 30% of the sequences analyzed, varying from incorrect N and C termini prediction to errors in intron/exon boundary prediction and missing exons; the accession numbers for those genes are listed in *italics* in Table 1. These errors became evident because a number of sequence tags obtained by MS/MS did not match or only partially matched the predicted protein sequences and by cross-correlating theoretical mass and pI values with the experimental values determined from the 2-D gels (see below). By systematic matching of ESTs against the genomic sequences, in many cases we arrived at the correct protein sequence (see also Mann and Pandey, 2001). We have selected several examples to demonstrate some of these annotation problems and reannotation strategies and to discuss implications for identification and localization (Figures 5A to 5D).

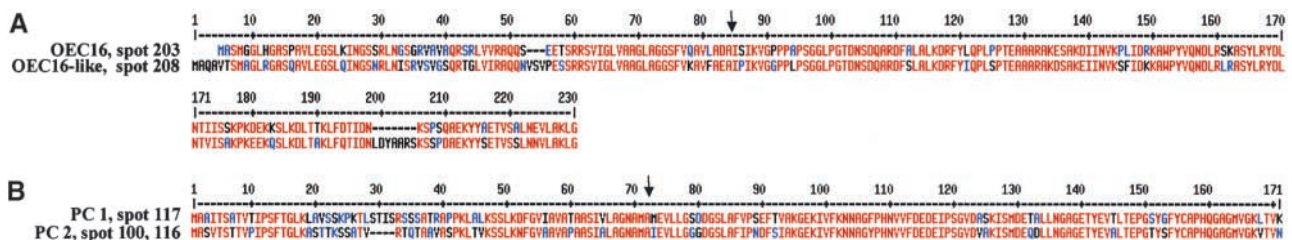


Figure 3. Sequence Analysis of Pairs of Isoforms/Paralogs Present in the Thylakoid Lumen or Associated with the Thylakoid Membrane of Arabidopsis Chloroplasts.

Alignments of two pairs of paralogs of thylakoid proteins identified on the 2-D gels shown in Figure 1. The cleavage sites of the ITP are indicated by the arrows.

(A) OEC16 isoforms in spots 203 and 208.

(B) Plastocyanin (PC) isoforms in spots 116 and 117.

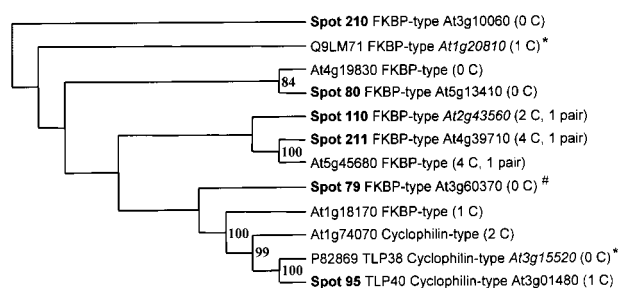


Figure 4. The Phylogenetic Relationship between Members of the Lumenal Isomerase Family.

The rooted tree of lumenal isomerases in spots 79, 80, 95, 110, 210, and 211, four predicted isomerases, and two isomerases found in Swiss-Prot (marked with *). The tree was built in Phylip using both parsimony and distance methods on protein sequences and reflects consensus trees with branches supported by the highest possible bootstrap values. The rooting was done with an outgroup sequence from the moss *Physcomitrella* and is an ortholog of spot 210. The location of Spot 79 is ambiguous (#).

The first example is a protein (spot 71, an OEC23 paralog) that was not annotated at all (Figure 5A). No ESTs for Arabidopsis were found either. We did not identify the protein by MALDI-TOF MS. However, we did identify two nucleotide stretches that matched the same region of chromosome II of the Arabidopsis genome by BLAST searching using two sequence tags determined by nano-ESI/MS/MS. The sequence tags also matched ESTs from soybean, and a protein of 214 amino acid residues could be reconstructed from overlapping EST sequences in soybean and potato. Alignment of these reconstructed genes with the nonannotated Arabidopsis genome sequence helped to identify the gene on chromosome II, and one additional sequence tag could be matched (Figure 5A). Directly N terminal of each sequence tag is a lysine or arginine, which confirms the reconstructed protein, because the protein was digested originally with trypsin, which cleaves C terminal of lysine or arginine. The Arabidopsis protein is predicted to contain a typical N-terminal chloroplast transit peptide and also has a putative ITP.

A second example is a peripheral fibrillin (spot 83) for which ~50% of the predicted protein was misassigned (Figure 5B). No ESTs are available for this protein either, but several ESTs were found for a closely related paralog. Comparison with this homologous fibrillin in Arabidopsis indicated that 240 bp of the sequence annotated as encoding sequence was likely to consist of introns and should be removed, whereas 309 bp was predicted to consist of introns but in fact consisted of exons. After correction of the fibrillin protein sequence, three sequence tags obtained by MS/MS matched to the sequence (Figure 5B).

The annotated genome sequence of a FK506 binding protein (FKBP) identified in spot 110 has a truncated N terminus (probably 60 to 70 amino acid residues are missing) starting

with MLLVL. Orthologs in tomato, barley, alfalfa, and the green alga *Chlamydomonas reinhardtii* all show bipartite presequences with typical lumenal twin-arginine motifs. The N terminus of the protein in spot 110 could be extended with 37 amino acid residues using one overlapping EST; however, no EST was found for the very N-terminal end. Several other proteins were found in which the N terminus was assigned incorrectly, leading to an incorrect localization prediction (see also Peltier et al., 2001).

More than 50% of the identified lumenal proteins have a typical twin-arginine motif, which is indicative of targeting through the TAT pathway. In many cases, all orthologs in other plant species also contain this TAT motif, indicating that the targeting route is conserved generally from algae to higher plants (although perhaps less conserved from cyanobacteria to plants [e.g., Hcf136] [Hynds et al., 2000]). However, spot 104 (Table 1) is a protein that has been shown experimentally to translocate via the TAT pathway (Mant et al., 1999). Database searching identified orthologs in seven other plant species. Interestingly, in five of these orthologs, the twin arginines were altered to KR, although the other features of a TAT motif were preserved. This finding suggests that in some cases TAT substrates also contain KR. Interestingly, very recently, a similar observation was made for an *Escherichia coli* protein (Hinsley et al., 2001). Further experimentation is needed to confirm that the orthologs with KR in fact translocate via the TAT pathway.

Correlation between Experimental and Theoretical Molecular Mass and pI Values, Interspecies Comparison, and Expression Levels

When searching the database with MS data, it is possible to use the pI and molecular mass values from the 2-D gels to reduce the number of proteins searched in the database. However, all lumenal localized proteins encoded by nuclear genes are synthesized with a bipartite presequence, and these presequences must be taken into account. To evaluate the contribution of the presequences of the lumenal proteins on pI and molecular mass, we cross-correlated the theoretical molecular mass and pI values of the precursors to those of the corresponding mature lumenal proteins (Figures 6A and 6B). The cross-correlation plot for the molecular mass shows a straight line parallel to a perfect correlation, with a shift of ~9 kD, and thus an average length of the cTP + ITP of ~81 amino acid residues (Figure 6A). This also indicates that the length of the presequence is independent of the size of the mature protein, which is very important for genome-wide prediction of the lumenal proteome (see below). Cross-correlation between the theoretical pI of the precursor versus the mature protein shows that the precursors are far more basic than are the mature proteins. Processing of the precursor proteins consistently gave a pI shift downward of up to 4 pI units, with generally larger shifts for the smaller proteins.

Table 3. Arabidopsis Proteins Copurified with the Thylakoid Membranes and Identified from the 2-D Electrophoresis Gels Shown in Figure 1

							Localization and Cleavage Site Prediction ^e		
Spot No.	Apparent Molecular Mass (kD)	pI	Identity ^a	Accession No. ^b	MALDI-TOF Percent Coverage at 50 ppm ^c	MS/MS Sequence ^d	TargetP	Predotar	SignalP
75	18.7	5.5	CipS1	5123926 (At4g25370)	35	AIAWAIDEK (1)	C1/63	P 1	−73 PIA-QP
92	33.3	6.2	Enoyl-acyl carrier protein reductase	4006834 (At2g05990)		C2/74	M 0.8	None	
111	19.5	6.5	Fe-superoxide dismutase	1351082 (At4g25100)	13	TFMTNLVSWEAVSAR (2)	No chloroplast	Neither	None
72	18.3	6.1	ROC4 isomerase	461899 (At5g13120)	21	FFMISYLDDEILIVR (6)	C2/67	P 0.8	−55-HYA-SP +47-GIA-LS
81	28.5	4.7	Fibrillin homolog	7484966 (At4g22240)	42		C2/59	P 1	None
82	28.9	4.7	Probable fibrillin	7488105 (At4g04020)	42		C2/55	P 1	None
83	28.2	4.9	Fibrillin CDSP34 homolog 1 +	6729544 (At3g58010)	40		C1/53	P 1	None
			Fibrillin CDSP34 homolog 2	2673904 (At2g42130)			LKEEYVEGMLETPTVIEEAVPEQLK (4)	C1/48	P 1
84	26.1	5.4	Similar to fibrillin	11994325 (At3g23400)	30	LIPVTLGQVFQR (3)	C1/72	P 0.8	−18 ALL-SD +104 LVA-SV
96	40.9	4.1	Putative fibrillin	3608139 (At2g35490)		C1/53	M 0.9	−45 YRP-KP +50 RFS-KI	
88	29.9	5.1	CP29 A' RNA-BP + Putative RNA-BP	681904 (At3g53460)	21	ILVDYLQDK (4)	C1/65	Neither	−12 AFN-PK
				3608147 (At2g35410)	13		C1/74	P 1	−45 SNL-SP +75 TSA-DE
119	32.1	4.1	CP31 RNA-BP	681908 (At4g24770)	40	35	C1/93	P 1	−85 DWA-EE +16 AMA-DS
85	24.3	4.8	CP29 B' RNA-BP	4056477 (At2g37220)	23	C1/47	P 1	−37 LSF-KL +107 AQL-AQ	
113	20.6	6.2	Similar to RNA-BP	4678944 (At3g52150)		21	C1/56	P 0.6	−42 SLA-GT
86	26.5	6.6	Ribosomal S5 P	4836870 (At1g78630)	20	41	C1/49	P 1	−17 LHT-RT +50 VKA-SS
107	17.1	5.3	Ribosomal S7 P	7525079	34		Chloroplast encoded	P 1	−20 LFL-SS +25 SHQ-IP
87	26.8	6.8	Ribosomal L4 P	3914666 (At1g07320)	41	29			
67	16.3	4.8	Ribosomal L12 P	548655 (At3g27850)	23	47	C1/54	P 0.8	−21 TCA-ST +58 VEA-PE
78	19.5	4.9	Ribosomal L13 P	1707008 (At2g33800)			C1/56	P 1	−19 VKS-SG +47 IYA-NS
65	15.9	4.7	Ribosomal L27 P	9759141 (At5g40950)	29	31	C4/19	P 1	−27 SFL-NR
64	15.2	4.9	Putative ribosomal recycling factor	7523401 (At3g63190)	40	C2/18	P 0.9	+48 LIA-CS	
209	18.9	9.3	Psa-D	4587564 (At1g03130)		40	C1/43	P 1	+43 AIR-AE
202	14.2	9.3	Psa-E	7443149 (At4g28750)	47	35	C2/44	P 1	±22 AGA-SS
38	33.9	5.6	FNR	5730139 (At5g66190)	21		C2/64	P 0.7	±48 VKA-QV
102	15.2	6.4	RbcS (20.2)	4204274 (At1g67090)	13	57	C3/54	P 1	±18 AQA-TM
129	50.6	6.6	RbcL (47.3)	1944432	41		Chloroplast encoded	P 0.9	−18 SLS-AD +43 SRA-SS
48 to 52	53.4	5.4 to 5.6	CF ₁ α (55.1)	5881679	57	22			
43 to 47	51.3	5.7 to 6.0	CF ₁ β (53.1)	5881701	57	Chloroplast encoded	P 0.8	−58 AMA-LA	
34	34.5	6.1	CF ₁ γ (53.1)	461550 (At4g04640)	22				53
8	17.8	5.8	CF ₁ δ (53.1)	5916447 (At4g09650)	55	Chloroplast encoded			
102 to 53	15.2	6.3 to 6.4	CF ₁ ε (53.1)	7525039	53				

These proteins are located at the stromal side of the thylakoids and were released by Yeda press. Proteins were identified by MALDI-TOF MS and/or nano-ESI/MS/MS.

^aIdentity given in NCBI completed by domain prediction found by Pfam, Blocks, Prints, Prodom, or Phi-Blast.

^bAccession numbers in NCBI and in MIPS (in parentheses).

^cPercentage of coverage at 50 ppm for the MALDI-TOF peptides.

^dSequence tag obtained identified by ESI/MS/MS; other sequences were obtained (number of sequences identified in parentheses) but are not shown.

^eLocalization, reliability class, and cleavage site prediction by TargetP (C for chloroplast), localization and score by Predotar (P for plastid, M for mitochondria), cleavage site prediction of the luminal transit peptide by two versions of SignalP developed for Gram-negative (-) or Gram-positive (+) bacteria.

We also cross-correlated the theoretical sequences (from the original annotation and reannotation) and experimental values for pI and molecular mass of the luminal proteins (Figures 6C and 6D). We, and many others for nonplant species, have observed that experimental pI and molecular mass generally correspond to the theoretical values. How-

ever, if there is a significant mistake in gene assignment, or if there is a large post-translational modification or processing event, these data points will lie outside of the correlation and thus will highlight such modifications or annotation errors. Generally, the correlation between predicted and experimental mass on the Arabidopsis maps were within a

Table 4. Small Set of Arabidopsis Proteins from the 2-D Electrophoresis Gel Shown in Figure 1 That Are Highly Abundant in Plant Cells and Are Not Localized in the Chloroplast

Spot No.	Molecular Mass (kD)	pI	Identity ^a	Accession No. ^b	MALDI Percent Coverage at 50 ppm ^c	MS/MS Sequence ^d	Localization
66	16.1	4.7	GCSH protein	121075 (At2g35370)	17	FFMISYLDDEILVR (2)	Mitochondria
97	40.1	6.7	Formate dehydrogenase	6625953 (At5g14780)	31		Mitochondria
98	59.4	4.7	Similar to PDI	5263328 (At1g21750)	42	TNVEVDQIESWVK (3)	Endoplasmic reticulum
94	36.7	5.8	Annexin	4959106 (At1g35720)		LLVSLVTSYR (2)	Cytosol
93	35.6	6.5	Annexin	4959108 (At5g65020)		LLLPLVSTFR (2)	Cytosol
76	22.0	6.8	GST	3201613 (At2g30860)	47		Cytosol
74	18.4	4.7	TCTP homolog	11994618 (At3g16640)		VVDIVDTFR (1)	Cytosol/nucleus

The cellular location for each protein is listed. Proteins were identified by MALDI-TOF MS and/or nano-ESI/MS/MS.

^aIdentity given in NCBI completed by domain prediction found by Pfam, Blocks, Prints, Prodom, or Phi-Blast.

^bAccession numbers in NCBI and in MIPS (in parentheses).

^cPercentage of coverage at 50 ppm for the MALDI-TOF peptides.

^dSequence tags identified by ESI/MS/MS and the number of sequences identified (in parentheses).

range of ± 5 kD (Figure 6C). However, three proteins (two PC spots and 204) were completely outside of this range and could correspond to the presence of homodimers. This dimerization could have occurred during the isoelectric focusing when the proteins reached their pI. The dimers apparently could not be solubilized before separation in the second dimension. Most of the pI values observed on the 2-D maps agreed with the corresponding theoretical pI values within a range of ± 0.5 pH unit. Four protein spots were in the range ± 0.5 to 1 pH unit. This larger difference is caused by post-translational modifications, visible on the gel as trains of spots (e.g., OEC23 spots 11 to 14), and might be caused by carbamylation. There are a number of post-translational modifications described for stromal and thylakoid proteins (glycosylation and palmitoylation; see Peltier et al., 2000); however, it is unknown whether such modifications occur on the luminal proteins.

One spot (spot 103A) showed a very large difference (>3 pH units) between predicted and experimental pI value (Figure 6D). This immediately suggested a possible problem of gene annotation. Therefore, the annotated genomic sequence was verified by overlapping ESTs. A reannotated gene was constructed, and the predicted protein sequence is shown (Figure 7, sequence B). Complete coverage of the reannotated gene was obtained and showed that exon IV had not been recognized as an exon and that exon V was predicted too short and was frameshifted. The theoretical pI value of sequence B still did not match the experimental value (Figure 6D). Interestingly, one additional EST covering the C-terminal end was found and a second protein sequence could be predicted (Figure 7, sequence C). In this case, exon VI was shorter than in sequence B, and two additional exons (exons VII and VIII) were added, as in the origi-

nal annotation. The theoretical pI value of the processed protein now matched the experimental pI value (Figure 6D), and 10 matching peptide masses from MALDI-TOF MS were identified. Thus, it is highly likely that the correct protein sequence of spot 103 is sequence C. This is a clear case of alternative splicing, which occurs (in the case of sequence C) in the middle of exon VI. In other cases, reannotation affected the theoretical pI only moderately (spots 106 and 108) or hardly at all (Figure 6D).

2-D maps of different plant species have been, and will be, generated for different subproteomes. To obtain a better insight into the interspecies correlation of 2-D gel patterns, we compared the maps generated for Arabidopsis with those generated earlier for pea (see Figure 2 in Peltier et al., 2000). Superficially, the pea and Arabidopsis 2-D maps show similar overall patterns, both in terms of expression levels and as coordinates for the different proteins, although the maps cannot be overlaid. To express this more quantitatively, we cross-correlated the molecular mass and pI values of the two species (Figures 8A and 8B). Nearly all pairs of pea/Arabidopsis proteins matched within a range of ± 10 kD and ± 0.75 pH, with the few exceptions indicated in Figures 8A and 8B. Interestingly, a nearly perfect correlation was found for spot 103, indicating that the acidic form was expressed preferentially in both organisms (Figure 7). All luminal proteins identified on the pea maps also were identified on the Arabidopsis maps, with the exception of two spots, indicating that the luminal proteomes of the two species are rather similar and showing the reproducibility of our preparations and 2-D gels. Naturally, more proteins were identified in the current study as a result of the completion of Arabidopsis genome sequencing and because the identification of pea proteins necessarily was based mostly on homology.

Sequence Analysis of cTP and ITP of the Experimentally Identified Proteins and Their Orthologs

To obtain cutoff parameters and selection criteria for a genome-wide prediction of luminal proteins (presented be-

low), we analyzed and assembled information concerning the ITPs of all experimentally identified and N-terminally sequenced luminal proteins, both from our own work and from the literature. The N termini for 41 nonredundant luminal proteins (including membrane proteins with ITPs, such

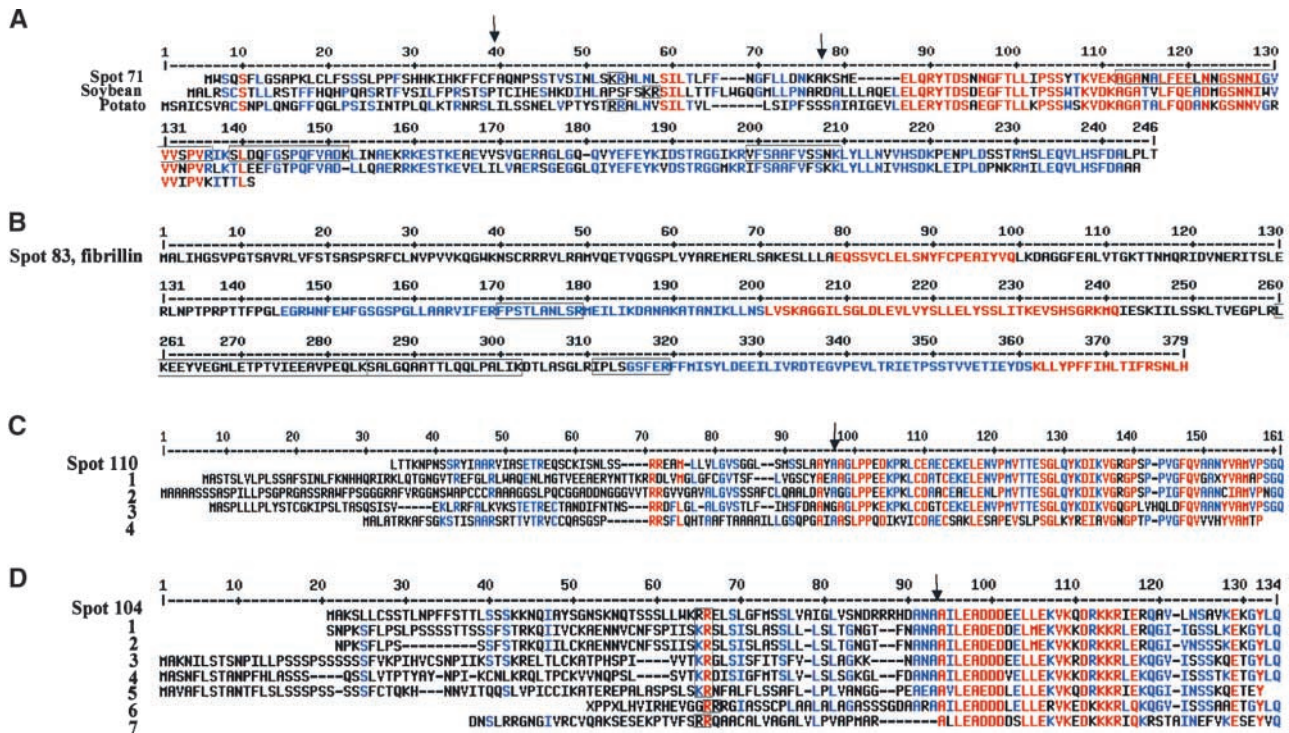


Figure 5. Correction of Gene Annotation and Other Sequence Analysis of Arabidopsis Proteins Identified on the 2-D Gels in Figure 1.

(A) Spot 71, identified as a 17.2-kD protein at pI 6.0 on the 2-D gel, was not annotated in the Arabidopsis genome, and no overlapping EST could be found. However, a homolog could be reconstructed in soybean using three overlapping ESTs. Using the reconstructed cDNA from soybean, the corresponding gene could be identified in the Arabidopsis genome on chromosome II. Three internal sequences determined by nano-ESI/MS/MS matched sequences in the gene and are indicated by boxes. Analysis with the functional domain predictor Pfam indicated that the protein in spot 71 belongs to the OEC23 family. The predicted ITP or cTP cleavage sites are indicated with arrows. Amino acid residues conserved among all three sequences are shown in red, and those conserved between two sequences are shown in blue. RR and KR motifs in the ITP are boxed.

(B) Spot 83 is a fibrillin (on chromosome II) and is an example of a serious misassignment in intron/exon boundaries. The misassignments were corrected by matching of the genomic sequence with a homologous fibrillin (verified entirely by overlapping ESTs) on chromosome III in Arabidopsis (T10K17.220) and verified by matching protein sequences obtained by nano-ESI/MS/MS; these sequence tags are boxed. To arrive at the correct amino acid sequence, amino acids shown in red need to be removed from the annotated sequence, and amino acids shown in blue need to be included. Fifty percent of the protein sequence was changed compared with the predicted sequence.

(C) The annotated genome sequence of a FKBP (accession number 22989010) identified in spot 110 has an incorrectly predicted N terminus (MLLV...). Orthologs in tomato (1) (AW041520), barley (2) (11193249), alfalfa (3) (11902372), and *C. reinhardtii* (4) (AV624465) all show typical bipartite presequences with typical luminal TAT motifs. The N terminus of the protein in spot 110 could be extended with one overlapping EST (T76027); however, no EST was found for the very N-terminal end. Amino acid residues conserved among all four sequences are shown in red, and those conserved between fewer than four sequences are shown in blue. The cleavage site for the ITP is indicated with an arrow.

(D) Alignment of the Arabidopsis protein sequence of spot 104 with its homologs in seven other plant species. Spot 104 has a typical TAT motif, and in vitro analysis has shown that this protein translocated via the TAT pathway (Mant et al., 1999). Interestingly, the first arginine residue of the twin arginines is replaced by a lysine residue in four of the orthologs. Orthologs are from potato (1) (10447846), tomato (2) (5900060), soybean (3) (7795408), alfalfa (4) (11900582), *Mesembryanthemum crystallinum* (5) (8330419), barley (6) (11198348), and *Physcomitrella patens* (7) (6102372). Amino acid residues conserved among all seven sequences are shown in red, and those conserved between fewer than seven sequences are shown in blue. The cleavage site for the ITP is indicated with an arrow.

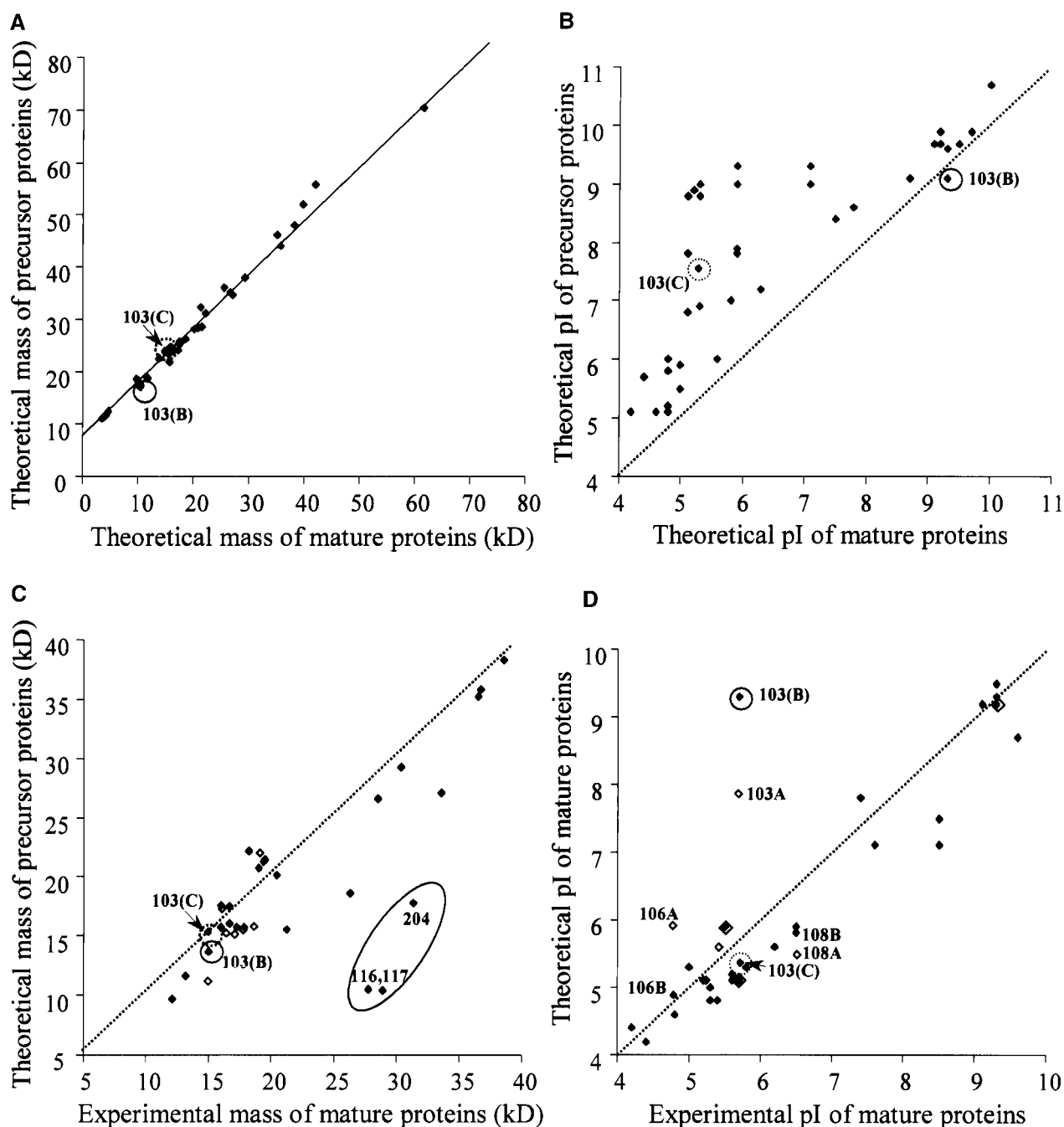


Figure 6. Cross-Correlation between Experimental and Theoretical Molecular Masses and pI Values of Precursors and Mature Proteins in Arabidopsis.

Cross-correlation of predicted and experimental molecular mass (**[A]** and **[C]**) and pI values (**[B]** and **[D]**) of the proteins from Arabidopsis identified in Figure 1 before (**[A]** and **[B]**) and after (**[C]** and **[D]**) removal of the cTP and ITP. Dotted lines indicate perfect correlations. Protein spot numbers are indicated for strongly deviated points. The circled data points correspond to spot 103 (see Figure 7). Open symbols represent values based on incorrect annotation (e.g., 103A, 106A, and 108A) (**[C]** and **[D]**). Three (most likely) monodimers (spots 116, 117, and 204) are indicated in **(C)**.

as PsbY1,2, PsbW, and PsbX) were identified experimentally in Arabidopsis or other plant species. To obtain a more statistically significant set, closely related orthologs of these nonredundant luminal proteins were identified by BLAST searching of all plant and algal (*C. reinhardtii*) protein and DNA sequence entries. Thus, a total of 201 luminal proteins were identified, and 109 of these have a typical TAT motif. Approximately 10% were orthologs in green algae. The cTPs of the algae are not well predicted by TargetP (see Discussion), but their ITPs were very similar to those of higher plants. It is important to note that ITPs show much less sequence conservation than does the mature protein sequence, between both paralogs and orthologs, thus reducing the redundancy in this ITP set. The ITPs then were analyzed using logoplots of the sequences aligned according to the experimentally identified N termini (Figures 9A and 9C) or to the TAT motif (Figure 9B). In the logoplots, the sequence alignment is represented by a sequence of stacked letters in which the total height of the stack at each position shows the amount of conserved information, whereas the relative height of each letter shows the relative abundance of the corresponding amino acid (Schneider and Stephens, 1990).

The comparison between the logoplots in Figures 9A and 9C shows that the N-terminal charged domains contain almost exclusively arginine in the TAT substrates, whereas in the Sec substrates the charged amino acid is preferentially lysine. In the Sec substrates, the hydrophobicity of the core domain is more pronounced and the C-terminal polar region is shorter, with a preference for a proline at position -4. No significant differences were found for positions -1 and -3, and the consensus sequence for the ITP cleavage site remains AxX for both substrates.

The logoplot in Figure 9B shows the alignment of the luminal TAT substrates according to the TAT motif and shows that the TAT motif in plants is RRxh(h/u) (where x is any residue, h is any hydrophobic residue, and u is any uncharged residue) and is more degenerated than it is in bacteria, which has a consensus motif of S/TRRxFLK (Berks et al., 2000). We did not observe any significant differences between ITPs of Arabidopsis proteins and those of other plants species or between ITPs of monocotyledonous versus dicotyledonous plants.

To define additional cutoff values for a theoretical genome-wide prediction of luminal proteins, the length distribution of cTPs and ITPs was analyzed (Figure 9D). A total of 141 proteins were used for the length distribution of cTP and are part of the experimental training set of TargetP (Emanuelsson et al., 2000); 174 luminal proteins, which are part of the set of 201 luminal proteins described above (some of the 201 proteins missed the very N terminus), were used for the length distribution of the total presequence of the luminal proteins (cTP + ITP). The two distributions are almost Gaussian, with an average size of 54 amino acids for cTP and 81 amino acids for cTP + ITP and a maximum length of 129 for cTP and 151 for cTP + ITP.

Genome-Wide Localization Predictions and Analysis

With the information summarized in Figures 9A to 9D, we sought to predict the luminal proteome in Arabidopsis. The set of Arabidopsis sequences for the theoretical analysis was obtained from Munich Information Center for Protein Sequences (MIPS) (see Methods) and contained 25,460 entries, including functional annotations (if any). The general prediction strategy is summarized in Figure 10A, and the prediction process can be followed through Table 5. To predict the potential luminal proteins, the protein sequences were first processed through the plant version of TargetP (Emanuelsson et al., 2000), which discriminates between proteins targeted to the chloroplast, the mitochondrion, and the secretory pathway (and as a fourth group, "other" destinations). TargetP also predicts the cleavage site of the cTP. A total of 3646 proteins were predicted to be localized to the chloroplast (using all reliability classes) and were processed further through SignalP 2.0 HMM version (SignalP-HMM; Nielsen et al., 1997, 1999), which discriminates between sequences containing a cleavable signal peptide, a noncleavable signal anchor, or neither. SignalP was designed originally to detect signal peptides for secretion in bacteria and eukaryotes, but ITPs show strong similarities to the secretory peptides (von Heijne et al., 1989). Evaluation of ITP prediction by SignalP on the set of experimentally identified luminal proteins listed in Table 1 shows no bias in successful prediction in either Gram-positive or Gram-negative mode. The performance of SignalP in the eukaryotic mode was less successful (Table 1). Thus, we used the union of the Gram-negative and Gram-positive versions of SignalP-HMM.

To mimic the actual sorting process, we removed the predicted cTP before submitting the sequences to the SignalP-HMM predictor (Figure 10A). Our first idea was to remove the stretch of residues corresponding to the TargetP-predicted cTP cleavage site, but that was abandoned because the TargetP prediction of cTP cleavage sites is quite inaccurate (~45% correct within two amino acid residues), in contrast to the more reliable chloroplast localization prediction (85% success rate). Instead, we processed the 3646 predicted chloroplast-targeted sequences by removing from 20 to 80 amino acids from their N termini in steps of five residues (90% of the cTPs in the TargetP training set fall into this length interval; Figure 9D) and submitted these sequences to SignalP for ITP prediction. Thus, for each predicted chloroplast protein, we obtained 26 SignalP predictions (in total, 13 processed versions of the sequences [removing 20, 25, 30, . . . 80 residues]) using both Gram-positive and Gram-negative versions of SignalP-HMM (Figure 10A). The prediction results are based on the union of these predictions and resulted in 1224 proteins with potential ITPs (Table 5, result 3). The sensitivity of the prediction at this stage was 94% (thus, there were 6% false negatives for the known luminal proteins), but, as expected, there was a strong overprediction on the set of 211 stromal proteins, resulting in a false positive rate of 38%.

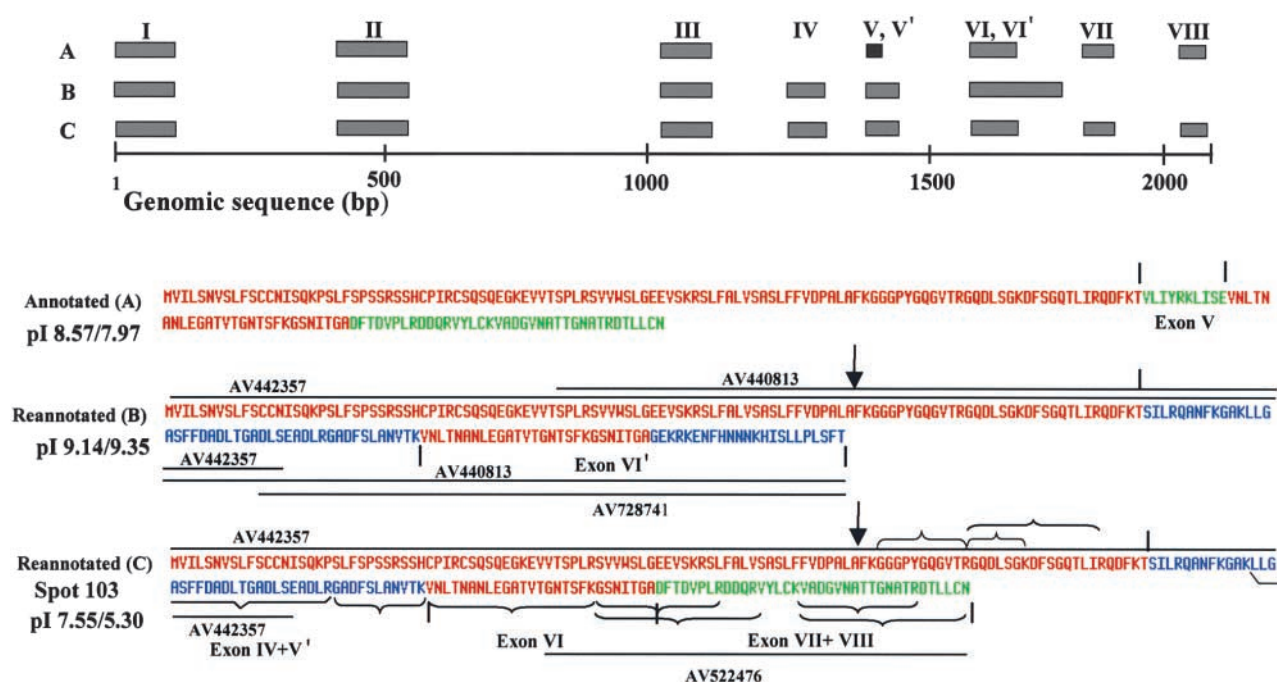


Figure 7. Alternative Splicing of Luminal Protein Spot 103.

The gene annotation for protein spot 103 is incorrect in the database. The top of the figure shows a scheme of the genomic sequence and the positions of the exons. Based on overlapping ESTs, it is clear that the gene annotation in the database (sequence A) is incorrect. Exon IV was not recognized as an exon, and exon V is too short and was frameshifted (indicated in black). One full-length cDNA and corresponding proteins (sequence B) can be reconstructed from the overlapping ESTs. Sequence B is constructed from six exons (I, II, III, IV, V, V', and VI) and encodes a protein with pI value of 9.35 for the precursor and 9.14 for the mature protein. Sequence C is constructed from exons I to V' plus exons VI, VII, and VIII and encodes a protein with a pI value of 7.55 for the precursor and 5.30 for the mature protein. The alternative splice site (in sequence C) occurs in the middle of exon VI. Two ESTs were used to reconstruct the protein in sequence C. The pI value and molecular mass of the processed protein match exactly the experimental coordinates on the 2-D gel (Figure 1A). Ten peptide masses determined by MALDI-TOF MS match this protein (mass accuracy within 50 ppm). Red indicates conservation, and blue or green indicates no conservation of the sequence between the different annotations A, B, and C. Matching Arabidopsis ESTs are indicated. The luminal processing site is indicated by the arrows.

Constraints then were added in the form of a demand for the $-3, -1$ motif to be present at the cleavage site, as shown in the logoplots of Figures 9A and 9C. The residues in the $-3, -1$ motif are preferentially alanines, but there also can be other small and apolar residues at the -3 position and (with very low frequency) at the -1 position (Figures 9A and 9C). The motif was applied in four versions of various stringency: (1) the restrictive (r) motif A-x-A (present at 75% of ITP cleavage sites in the set of 201 known luminal proteins); (2) the average (a) motif [ASV]-x-A (present in 87%); (3) the semipermissive (s) motif [ADGLSTV]-x-A (present in 95%); and (4) the permissive (p) motif [ADGLSTV]-x-[AGS] (present in 97%). The p motif included all amino acids that were present more than twice at positions -1 and -3 in the set of 201 luminal proteins. For each motif, a list of cTP + ITP-containing proteins was generated, and these are summarized in Table 5. This showed that the sensitivity (tested on the known luminal proteins) decreased from 0.94 to 0.59

with increasing stringency, whereas the false positive rate decreased from 0.38 to 0.17 (tested on the set of known stromal proteins).

Additionally, to be considered a potential luminal protein, the total length of the predicted cTP + ITP was required to be between 60 and 150 residues (corresponding to experimentally verified lengths in the set of 174 complete luminal proteins [Figure 9D]), and the processed mature part of the protein was required to be at least 50 amino acid residues. Using the p motif, 774 proteins were left in result 5, whereas applying the r motif reduced the set to 190 proteins (Figures 6A and 10A). At this stage, the sensitivity was 0.85 and 0.56 and the false positive rate was 0.21 and 0.1 for the p and r motifs, respectively.

Next, we verified the sequences for potential TM regions using the predictor program TMHMM (version 2.0) (Krogh et al., 2001). All sequences that were predicted to contain a TM region either overlapping with the ITP cleavage site or

downstream of it were discarded, because our aim was to find soluble luminal proteins. TM domains inside of the ITP regions were not considered because such proteins should have been predicted as signal anchors by SignalP-HMM. Some of the proteins that were discarded at this step are membrane proteins with an ITP, such as PsbW, PsbX, and PsbY1-Y2.

Among the remaining sequences, potential luminal proteins with the TAT motif R-R-xh(h/u) were collected (Figure 9B). The twin arginines in the set of 201 known luminal proteins are located 18 to 32 amino acid residues upstream of the ITP cleavage site. Consequently, we used the presence of two adjacent arginines, present within -18 to -32 relative to the ITP cleavage site, as a criterion. They also were required to be at least 35 residues downstream of the N terminus of the precursor protein. This resulted in sets of 27, 52, 62, and 93 TAT proteins, depending on the $-3, -1$ motif requirement. For the p motif, only 6% of the 211 known stromal proteins were falsely predicted as luminal with a TAT motif, and the sensitivity was 83% (Table 5). In the set of 93 proteins (p motif), only 30% were annotated functionally in MIPS. Therefore, all proteins were passed manually through all major functional domain predictors, increasing (often very limited) annotation to 56% of the proteins. Then, 22 proteins (24%) with known or predicted functions obviously incompatible with luminal localization were excluded; the resulting list of 71 proteins is shown in Table 6. These 22 false positives included mostly proteins involved in transcription and translation and some potential nonchloroplast proteins. Forty-four percent of the remaining 71 proteins lack functional annotation (Figure 10B), and determining their function might help to fully understand the functions of the luminal proteome.

The prediction for the set of potential luminal proteins without a TAT motif was not as good as that for the TAT substrates, and the false positive rate was between 0.14 (p) and 0.09 (r), with sensitivities of 0.60 (p) and 0.40 (r). Therefore, these results are only summarized in Table 5.

DISCUSSION

The thylakoid lumen in chloroplasts is still a poorly characterized compartment, and only a few dozen proteins have been identified. A number of functions of the luminal proteome are well known, such as water splitting by the OEC (involving OEC16, OEC23, and OEC33) of photosystem II, electron transport (e.g., involving PC), and regulation of the xanthophyll cycle (violaxanthin deepoxidase). A number of additional luminal proteins have been identified with known functional domains, such as an isomerase (TLP40), a carbonic anhydrase, a photosystem II "assembly factor" (Hcf136), a protease (DegP1), and the thylakoid-processing peptidase and the D1-processing peptidase (CtpA). In addition, recent systematic searches identified a number of lu-

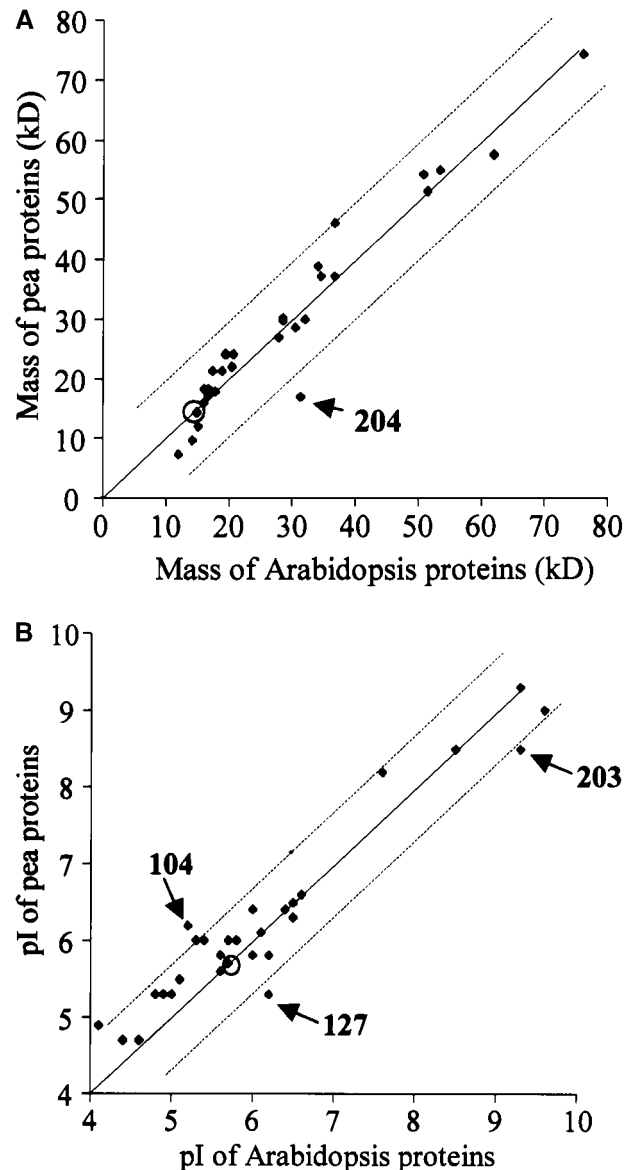


Figure 8. Cross-Correlation of the Experimental Molecular Mass (**A**) and pI Values (**B**) of the Luminal Proteins from Pea and Arabidopsis Identified on the 2-D Maps.

The broken lines indicate a deviation of 10 kD or 0.75 pI units. Protein spot numbers are indicated for strongly deviated points. The circled data points correspond to spot 103 (see Figure 7).

menal proteins of unknown function (Kieselbach et al., 1998; Peltier et al., 2000).

In this study, we sought to characterize the thylakoid lumen proteome further by a unique combination of experimentation and localization prediction. We also determined the relative expression levels, both to estimate the dynamic resolution of

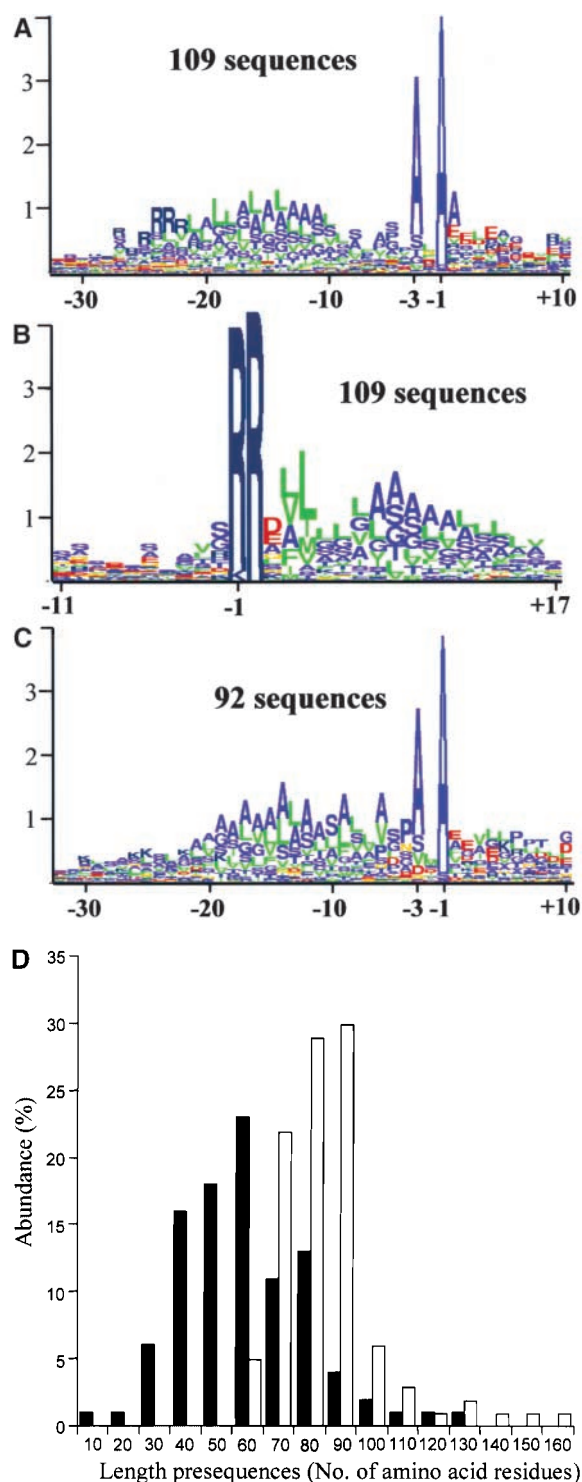


Figure 9. Sequence Analysis of cTP and ITP of the Experimentally Identified Proteins in Arabidopsis and Their Homologs/Orthologs in Other Plant Species.

(A) and (B) Logoplot of 109 luminal proteins with a typical TAT motif

our experimental setup and to obtain insight into stoichiometries between different proteins and their paralogs.

Copurification of Peripheral Proteins

In addition to the luminal proteins, we identified many proteins that are attached to the stromal side of the membrane as part of their function. It was natural that these peripheral proteins copurify with the luminal proteins because they are released easily by either Yeda press or the sonication used to open the luminal compartment. It is important to emphasize that few proteins were identified on the gels that can be considered true contaminants, such as the abundant proteins from the stroma or from outside of the chloroplasts. This is the result of multiple wash steps with buffers of low ionic strength and purification of the membranes on Percoll gradients. We did attempt to reduce the copurification of proteins located at the stromal side of the membrane by using different sonication times and energies. Overall, Yeda press treatment of the luminal proteome gave the “purest” luminal proteome (as judged by the low amounts of released $CF_1\alpha,\beta$ [~ 5 to 10% of the total]) while still obtaining good efficiency of extraction, as judged by the protein yield per milligram of chlorophyll and the amount of extracted membrane-bound OEC33. It is important to remember that the luminal proteins can be soluble or bound peripherally (e.g., OEC33) to the luminal side of the thylakoid membrane or both; thus, a minimal amount of energy is needed to release these proteins. We have considered protease treatments of thylakoids before extraction of luminal proteins (as is often used for localization of individual proteins in combination with protein gel blotting), but this is likely to produce many breakdown products that will copurify with the lumen, thereby making the maps irreproducible and the identification process very difficult.

Thus, the maps in the study represent the soluble luminal proteins and weakly, peripherally attached proteins at either side of the thylakoid membrane. Proteins that interact more tightly with the membrane, either through TM domains or lipid anchors, are not present on the 2-D gels.

identified experimentally in Arabidopsis and their homologs in other plant species aligned according to the experimentally identified N terminus **(A)** and RR motif **(B)**.

(C) Logoplot of 92 luminal proteins without a typical TAT motif identified experimentally in Arabidopsis and their orthologs in other plant species aligned according to the experimentally identified N terminus. **(D)** Length distribution of cTP for the experimental training set of TargetP (141 proteins) (closed bars) and the length distribution of cTP + ITP (open bars) for the 174 experimentally identified luminal proteins. Proteins are divided into classes of 10 amino acid residues.

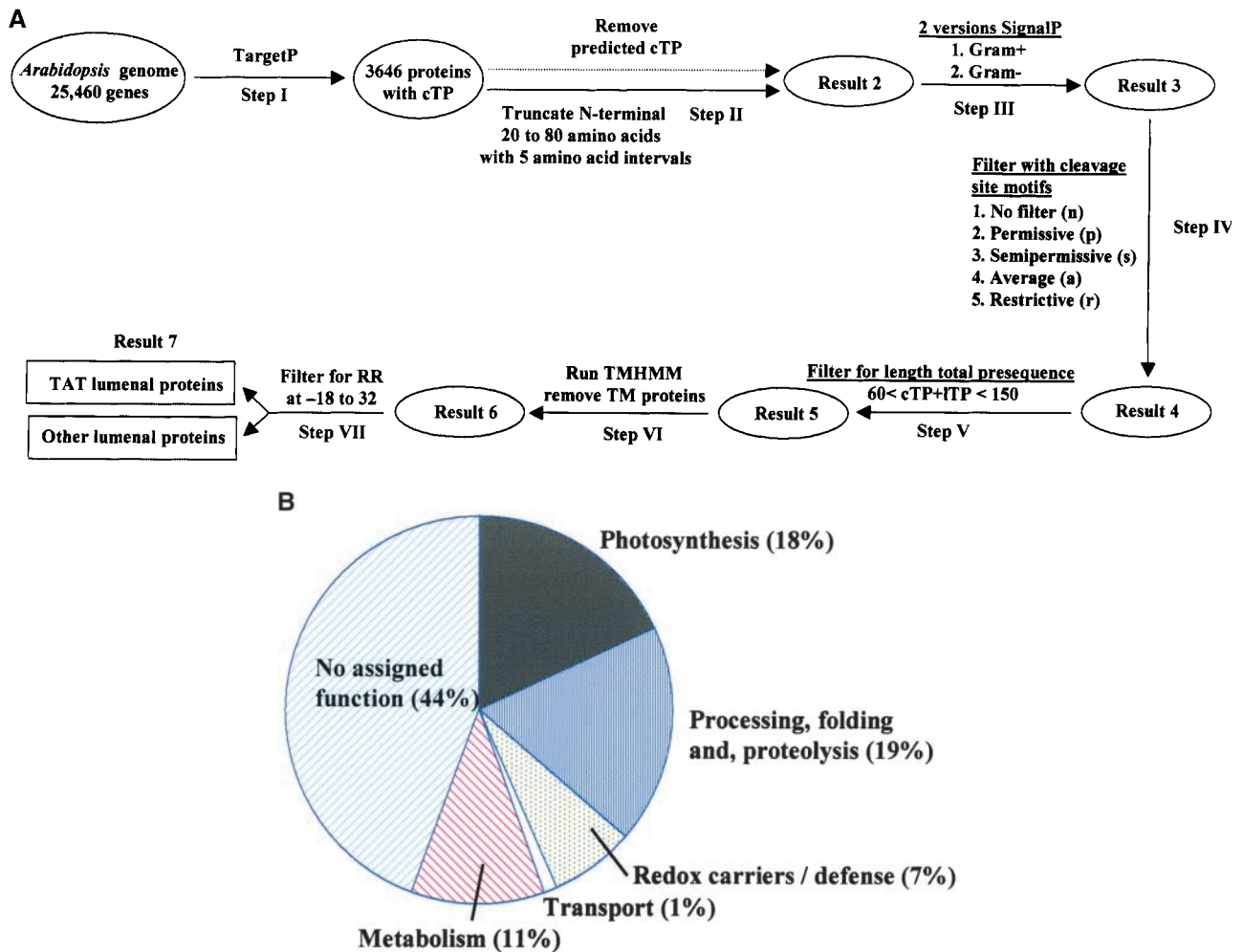


Figure 10. Genome-Wide Prediction of the Luminal Proteins with a Typical TAT Motif in the Genome of Arabidopsis.

(A) Summarizing scheme of the genome-wide prediction of luminal proteins with a TAT signal.

Step I. 25,460 Arabidopsis open reading frames (see Methods) were processed through TargetP, resulting in 3646 protein sequences predicted to have a cTP (Result 1).

Step II. For each of the 3646 proteins, 20, 25, . . . 80 residues were removed from the N terminus to mimic the cleavage of the cTP. (Thus, each protein was present in 13 differently truncated versions.)

Step III. The truncated proteins were processed through SignalP (both Gram-negative and Gram-positive versions) to predict the potential presence of a ITP. If any of the 13 truncated versions of a protein were predicted to contain a ITP by at least one of the two SignalP versions, the protein was kept in Result 3 (1224 proteins).

Step IV. The proteins predicted to have a ITP were checked for the presence of the four versions of the $-3, -1$ motif (p, s, a, and r; see Results) at the ITP cleavage site.

Step V. Length restrictions were imposed. Proteins that did not meet the length criteria were excluded (Result 5; 596 proteins using the p motif). Step VI. The remaining proteins were processed through the TM region predictor TMHMM. All proteins that contained one or more TM regions in the predicted mature part (i.e., C terminal of the predicted ITP cleavage site) were removed.

Step VII. Proteins with a TAT pathway motif (twin arginine; RR) in region -32 to -18 relative to the predicted ITP cleavage site were sorted to the TAT luminal protein set (93 proteins using ITP cleavage site motif p). The rest of the proteins were kept in the "other luminal proteins" set (380 proteins using the p motif).

(B) Functional catalog of the 71 predicted luminal proteins with a TAT motif based on known function or functional domain prediction.

Table 5. Summary of the Different Result Steps during the Genome-Wide Screening for Lumenal Proteins, following the Strategy Shown in Figure 10A

Result Steps (see Figure 10A)	No. of ITPs	False Positive Rate on the Set of 211 Known Stromal Proteins	True Positive Rate on the Set of Experimentally Identified Lumenal Proteins (Sensitivity)
Result 3	1224	0.38	0.94
Result 4			
n	1224	0.38	0.94
p	943	0.34	0.94
s	630	0.29	0.88
a	525	0.27	0.76
r	230	0.17	0.59
Result 5			
p	596	0.29	0.85
s	411	0.18	0.82
a	343	0.17	0.74
r	161	0.12	0.56
Result 7, TAT			
p	93	0.06	0.83
s	62	0.05	0.83
a	52	0.05	0.67
r	27	0.02	0.39
Result 7, other			
p	380	0.14	0.60
s	267	0.12	0.53
a	221	0.11	0.47
r	105	0.09	0.40

Expression of Known Lumenal Proteins

From the 2-D gels of Arabidopsis, we identified 30 lumenal proteins and an additional 12 potential lumenal proteins. The expression levels of these identified proteins varied greatly, with a molar range of 1 to 10,000 (Figure 2). However, using narrow-gradient gel multidimensional chromatography combined with on-line nano-ESI/MS/MS measurements, it is technically possible to identify these proteins of lower abundance and reach a dynamic resolution of 1:100,000 molar ratio or better.

We identified all of the very abundant known lumenal proteins involved in photosynthesis, such as OEC16, OEC23, OEC33, PC, and PsaN, with relative expression levels of 50 to 100 compared with Hcf136. We should note that the expression levels of these abundant proteins were underestimated by a factor 3 to 5 because of saturation of the stains. We also identified the abundant nonphotosynthetic lumenal proteins described in earlier studies, such as Hcf136, TLP40, and TL16.5, with relative expression levels of ~ 1 to 10.

Expression of Paralogs of Photosynthetic Proteins and a New Putative Photosystem II Protein

A closely related paralog was identified for OEC33, PC, and an OEC16 at a relatively high expression level (5 to 50). The

pairs of PC and the OEC16 paralogs were well conserved at the primary sequence level (with 83 and 73% identity of the mature protein, respectively) (Figures 3A and 3B); thus, it is possible that they do function in photosynthesis. However, it is unclear why both members of each pair are expressed.

In addition, we identified seven paralogs of OEC23 on the Arabidopsis maps, and an eighth was identified earlier on our pea maps (spot 22). A ninth member was identified in the database and was 88% identical as OEC23, but it is unlikely to be expressed. The group of seven expressed OEC paralogs is relatively distant from OEC23 (and its nonexpressed homolog), with 18 to 31% identity; therefore, it is quite likely that they do not function in the OEC, which leaves their function completely unknown. Expression levels were 20 to 300 times lower than that of OEC23.

The presence of the paralogs of OEC16, OEC23, OEC33, PC, and others indicates gene duplication. Attempts to date these gene duplication events suggested that some of these duplications occurred around the time of the divergence of *Brassica* and the Rosidae (data not shown).

A very basic (pI 9.3) lumenal protein without predicted TM domains and with homology with a cyanobacterial 11-kD protein in photosystem II was identified in both pea and Arabidopsis. We tentatively assigned it as a new photosystem II protein, but its precise localization in the lumen and its function remain to be determined.

Expression of Multiple Isomerases and Proteases

Isomerases belong to two large families of proteins: the protein disulfide isomerases (PDI) and the peptidyl-prolyl *cis-trans* isomerases or rotamases. In addition to thiol-disulfide exchange, PDIs catalyze the GSH-dependent reduction of dehydroascorbate, bind ions and hormones, serve as a subunit of prolyl hydroxylase, and show triglyceride transferase activity (Gilbert, 1998). The PDI-like activity is mediated either by the thioredoxin-like CXXC motif with two vicinal cysteines or by a single, highly reactive cysteine; in the latter case, GSH can interact with the mixed disulfide instead of the second cysteine and release the substrate protein (Walker and Gilbert, 1997). The peptidyl-prolyl *cis-trans* isomerases or rotamases, which catalyze the *cis-trans* isomerization of the amide bond between a proline residue and the preceding amino acid residue, are composed of at least three different groups: the cyclophilins inhibited specifically by cyclosporins, the FKBP's inhibited by FK506 and rapamycin, and the parvulins inhibited by juglone (Schiene and Fischer, 2000).

Five isomerases were identified in the lumen (Table 1), and one is listed in Table 2. They can function in folding of the thylakoid proteins or in signaling (such as TLP40) or possibly are part of the antioxidant defense system via a connection to 2-Cys peroxiredoxins (see next paragraph). Only TLP40 has been described in more detail (Fulgosi et al., 1998; Vener et al., 1999): it is characterized by a cyclophilin-like C-terminal segment of 20 kD, a predicted N-terminal leucine zipper, and a potential phosphatase binding domain (Fulgosi et al., 1998). The isolated protein possesses peptidyl-prolyl *cis-trans* isomerase protein folding activity, and TLP40 also exerts an effect on the dephosphorylation of several key proteins of photosystem II. The genome-wide prediction of luminal proteins identified four additional chloroplast isomerases with potential ITP with a TAT motif. One additional isomerase with a TAT motif and an isomerase targeted via the Sec (or other) pathway were found with Swiss-Prot, increasing the set of potential luminal isomerases to 12. The relationships in this isomerase family are shown in Figure 4 in the form of a rooted tree with three main clades. None of these isomerases were identical to other known chloroplast isomerases, such as FKBP13 (Luan et al., 1994a), ROC4 in the stroma (Lippuner et al., 1994), cyclophilin pCyP B (Luan et al., 1994b), RB60, a protein disulfide isomerase involved in translation regulation (Kim and Mayfield, 1997; Trebitsh et al., 2000), or any of the other known cytosolic isomerases (ROC1, -2, -3, -5, or -6) (Chou and Gasser, 1997). On the basis of the presence of a closely spaced cysteine pair, it is possible that spots 110 and 211 and At5g45680 (indicated in Figure 4) have a specific function as an oxidative folding catalyst. These three isomerases also are clustered. Five of the isomerases have no cysteines at all and likely function as prolyl *cis-trans* isomerases only, whereas the four remaining isomerases have one or two cysteines and possibly play a role in oxidation/reduction reactions.

Antioxidant Metabolism in the Thylakoid: A Network of m-Type Thioredoxins, Peroxiredoxins, and Ascorbate Peroxidase

Light-driven charge separations in the photosystems and side reactions in photosynthetic electron transport such as the Mehler reaction, as well as photorespiration in the peroxisomes, produce reactive oxygen species that need to be quenched to prevent extensive damage to proteins as well as membrane lipids. Different carotenoids present in the photosystems and their antennae, as well as a network of superoxide dismutases and pools of vitamin E, ascorbate, and GSH, form a strong antioxidant network in and around the thylakoid membrane (Noctor et al., 2000). However, it is not clear how much this network extends into the thylakoid lumen.

The luminal protein violaxanthin deepoxidase is involved in the xanthophyll cycle and uses ascorbate as a cofactor, implying directly that ascorbate must be present in the lumen. However, no evidence for ascorbate in the lumen has been shown, which might relate to experimental difficulties of its identification rather than to its absence from the lumen. It is not clear how ascorbate is transported into the lumen, and neither is it clear if oxidized ascorbate can be rereduced in the lumen (either spontaneously or by enzymatic reaction) or if it is exported in its oxidized form (as monodehydroascorbate or dehydroascorbate) back into the stroma, where it is reduced by monodehydroascorbate reductase or dehydroascorbate reductase.

On the 2-D map, we identified a putative ascorbate peroxidase with all of the characteristics of a luminal protein and distinct from the known ascorbate peroxidases soluble in the stroma and one bound to the thylakoid membrane (Jespersen et al., 1997; Yoshimura et al., 1999) and identified earlier on our 2-D gels from pea. Its precursor has a typical ITP with a TAT motif, and the experimentally determined N terminus corresponded with the prediction by SignalP, suggesting that this ascorbate peroxidase is in fact located in the thylakoid lumen.

A number of additional members of the antioxidative defense system were identified on the 2-D maps. They are a known thylakoid-located 2 Cys peroxiredoxin (2-CP) (Baier et al., 2000), an uncharacterized peroxiredoxin-like protein, and three m-type thioredoxins (m1, m2, and m4) functionally distinct from the f-type thioredoxins involved in the activation of Calvin cycle enzymes (Issakidis-Bourguet et al., 2001). 2-CPs are peroxidases that detoxify alkyl hydroperoxides (resulting from lipid oxidation) by reduction to their corresponding alcohols. Antisense plants with reduced 2-CP levels showed severe damage to the photosynthetic apparatus and upregulation specifically of the known ascorbate peroxidases in the stroma and thylakoid, without much effect on GSH metabolism, superoxide dismutases, or catalase (Baier and Dietz, 1999; Baier et al., 2000). Interestingly, the peroxiredoxins link both to the ascorbate pathway and to the cyclophilins, because it has been shown in mammalian

Table 6. Seventy-One Predicted Luminal Proteins with a TAT Motif, According to the Scheme Shown in Figure 10A, Using a Permissive Cleavage Site of the ITP^a

Name	Length (eTP ITP)	Position of RR from N Terminus	ITP Cleavage Site Motif (s/p)	Functional Annotation (According to MIPS)	Additional Functional Annotation (This Article) and Match to 2-D Gels
At1g03600	68	41	a	Unknown protein	Spot 207, putative new photosystem II protein
At1g05420	86	55	a	Hypothetical protein	
At1g06430	73	48	r	Cell division protease Ftsh, putative	
At1g06680	77	55	r	23-kD polypeptide of OEC	
At1g14150	65	43	r	Unknown protein	Homology with OEC16
At1g15510	64	36	p	Hypothetical protein	Pentatricopeptide repeat and prokaryotic membrane lipoprotein attachment
At1g17210	64	36	a	Hypothetical protein	
At1g18170	94	74	p	Hypothetical protein	Putative FKBP isomerase
At1g21500	59	41	r	Unknown protein	
At1g26360	62	34	p	Hypothetical protein	Epoxide hydrolase signature
At1g35210	90	62	p	Hypothetical protein	
At1g49630	90	61	p	Hydrogenase protein, putative	
At1g55580	60	42	s	Hypothetical protein	
At1g70350	81	54	r	Hypothetical protein	
At1g71200	93	72	p	Hypothetical protein	Related to myc protein
At1g73530	73	46	p	Hypothetical protein	
At1g74070	74	45	a	Hypothetical protein	Putative cyclophilin
At1g76450	80	57	r	Unknown protein	Spot 70
At1g77090	63	47	s	Unknown	Spot 108, OEC23 related
At2g01400	85	63		Hypothetical protein	Related to Lon protease <i>Brevibacillus</i> 402504
At2g15570	68	45	a	Putative thioredoxin m	
At2g20270	62	39	a	Putative glutaredoxin	
At2g23670	71	49	a	Hypothetical protein	
At2g26340	80	59	r	Unknown protein	
At2g30790	76	56	a	Putative photosystem II OEC23 protein	See Results and Discussion
At2g34860	93	64	r	Unknown protein	Related to chaperone HSP40/DnaJ
At2g37240	68	45	s	Unknown protein	Related to protein 13834657 Mus
At2g37660	86	62	p	Unknown protein	Related to 3β HSD isomerase
At2g39080	72	40	p	Unknown protein	
At2g39470	73	51	s	Unknown protein	Spot 212, OEC23 related
At2g40400	93	67	r	Unknown protein	Related to protein 7572912
At2g47390	98	51	s	Unknown protein	
At3g01440	74	50	a	Hypothetical protein	Related to OEC16
At3g03760	77	59	p	Unknown protein	Related to 13569546
At3g05020	66	42	a	Acyl carrier protein 1 precursor	
At3g10060	82	60	r	Unknown protein	Spot 210, putative isomerase
At3g10130	83	49	s	Unknown protein	Related to 13877685 and 13424114 from <i>Caulobacter</i>
At3g11630	69	51	a	Putative 2-Cys peroxiredoxin	Spot 74 (Table 2)
At3g16000	95	73	r	Myosin heavy chain-like protein	
At3g52960	71	52	a	Peroxioredoxin-like protein	Spot 68 (Table 2)
At3g55330	74	50	a	Putative protein	Spot 213, OEC23 related
At3g56140	90	65	a	Putative protein	Related to 4586056
At3g57680	99	74	r	C-terminal protease-like protein	
At4g05180	82	60	a	Oxygen-evolving enhancer protein 3 precursor-like protein (OEC16)	Spot 208
At4g09010	82	53	r	Putative protein	Spot 205 to 206, putative ascorbate peroxidase
At4g15120	95	58	p	Hypothetical protein	Related to 11994735
At4g15510	104	79	r	Hypothetical protein	Spot 19, OEC23 related
At4g19830	78	52	r	Putative protein	Putative FKBP isomerase
At4g21280	75	53	a	Photosystem II OEC protein 3-like (OEC16)	Spot 203
At4g25130	71	44	r	Protein-methionine-S-oxide reductase	
At4g26500	75	48	p	Putative protein	protein Duf and BolA family
At4g29590	85	54	r	Putative protein	Related to methyltransferases
At4g31390	61	30	a	Predicted protein	Related to ABC transporter
At4g31560	75	45	s	Putative protein	
At4g32020	71	42	a	Putative protein	Related to 13272401
At4g34020	80	52	p	Putative protein	ThiJ family
At4g34120	72	42	a	Putative protein	CBS domain (protein interaction)
At4g36530	63	41	a	Putative protein	Epoxide hydrolase signature
At5g04900	63	36	a	Putative protein	
At5g11450	95	70	s	Putative protein	Spot 22, pea map OEC23 related
At5g11550	60	42	p	Putative protein	
At5g13410	86	61	s	Putative protein	Spot 80, putative FKBP isomerase
At5g17710	67	43	p	Chloroplast GrpE protein	Spot 89
At5g23120	78	55	r	Photosystem II stability/assembly factor Hcf136	Spot 123
At5g27860	71	52	p	Putative protein	
At5g39830	92	67	p	DegP protease-like protein	DegP8
At5g45680	79	58	r	Putative protein	Putative FKBP isomerase
At5g50110	66	40	p	Putative protein	
At5g55570	68	37	s	Unknown protein	
At5g62840	63	37	r	Putative protein	
At5g64040	86	63	s	Photosystem I reaction center subunit psaN precursor (PSI-N) (sp P49107)	Spot 201

^aFrom the 93 proteins originally predicted, 22 (24%) were removed manually because they were likely to be false positives. Matches to experimentally identified protein spots are noted.

tissue that cyclophilin A binds to peroxiredoxins, suggesting that cyclophilin A acts as an immediate donor of peroxiredoxins (Lee et al., 2001). If this is relevant for the chloroplast homologs, it would link several cyclophilins to different peroxiredoxins.

The N termini of m2 and m4 thioredoxins on our map were sequenced and confirmed chloroplast localization. Recently, four m-type thioredoxins in Arabidopsis have been implicated directly in antioxidant defense (Issakidis-Bourguet et al., 2001). By complementation assays with a mutant yeast strain, it was found that m1, m2, and m4 thioredoxin and one x-type thioredoxin, but not the f-type thioredoxins, induced hydrogen peroxide tolerance. Unexpectedly, the fourth m-type thioredoxin, m3, had a hypersensitizing effect on oxidative stress. Overall, the authors concluded that these m-type and x-type thioredoxins can serve as electron donors for the reduction of hydroperoxides (Issakidis-Bourguet et al., 2001). Possibly, these thioredoxins form direct redox couples with the peroxiredoxins, as was shown for Arabidopsis thioredoxin m3 with cytosolic thioredoxins (Verdoucq et al., 1999). It is interesting that we identified m1, m2, and m4 on our map but not m3, considering that only these three confer hydrogen peroxide tolerance in yeast. As is evident from the list of thioredoxins and peroxiredoxins in Table 2, we are not certain if these proteins are luminal proteins. Indeed, it seems that 2-CP is located at the stromal side of the membrane (K.J. Dietz, personal communication); the precise thylakoid localization of the other proteins remains to be determined.

Peripheral Proteins Involved in Translation, Stress Response, and Proteolysis

All but one of the set of 32 peripheral proteins (Table 3) were predicted by TargetP to be in the chloroplast (but not in the lumen) by the criteria of the prediction scheme in Figure 10A. A set of six (putative) fibrillins were identified, and some have been shown to be central in storing carotenoids of different plastids and accumulating plastoglobules (in chromoplasts, etioplasts, and chloroplasts) (Deruere et al., 1994; Rey et al., 2000). Overexpression and antisense suppression of different fibrillins affected development and stress response (Monte et al., 1999; Rey et al., 2000). Clearly, the fibrillins are sufficiently abundant to be identified on the 2-D gels, opening the way to study differential expression and the fibrillins' relationship to chloroplast development and stress in more detail. Because many proteins are translated at the thylakoid surface, it is not surprising to find a number of mRNA binding proteins (Table 3). Five mRNA binding proteins were identified, three of which have been described recently; however, their precise role is unclear (Nakamura et al., 2001).

Finally, we should focus attention on ClpS1 (Table 3). Recently, we identified ClpS1 as a member of a thylakoid-associated Clp protease complex of 350 kD with two (putative)

heptameric rings. ClpS1 shares weak homology with the ClpP proteins, and we suggested that ClpS1 is not part of the ring structure but is positioned at the axial opening of the ClpP/R core (Peltier et al., 2001). The identification of ClpS1 on the 2-D gel confirms the thylakoid association and suggests that it could function to link the large protease complex to the thylakoid surface, in agreement with a postulated role in the degradation of cytochrome *b₆f* subunits (Majeran et al., 2000).

Gene Annotation Errors and Generic Methods for Reannotation

For ~30% of the proteins listed in Table 1, significant errors in gene annotation were identified, varying from no annotation of the gene to mistakes in intron/exon boundaries, missed exons, and truncated N termini. These genes were reassigned using overlapping ESTs (if available) and verified by sequences obtained by MS/MS. To match the genomic sequences with overlapping ESTs, at least two World Wide Web sites are now available (<http://bioinformatics.iastate.edu/cgi-bin/gs.cgi> and http://www.tigr.org/docs/tigr-scripts/nhgi_scripts/), and it is expected that many of these misassignments will be corrected rapidly with overlapping ESTs if they are available. However, if no or limited matching ESTs are available for Arabidopsis, it will be very beneficial to reconstruct first one (or more) orthologous gene in another species and use that reconstructed gene to map it to the genomic Arabidopsis sequence, as was demonstrated in this study. To date, this interspecies matching can be done only manually (to our best of knowledge). Finally, we noticed discrepancies in gene annotation (intron/exon boundaries, etc.) between the MIPS and National Center for Biotechnology Information (NCBI) databases (e.g., spots 103, 106, and 211), but this will be resolved in coming years by The Arabidopsis Information Resource (TAIR) (S. Rhee, curator TAIR, personal communication).

TAT Proteins with a KR Motif?

BLAST searching with the known luminal proteins indicated that targeting pathways generally are conserved between orthologous proteins. In the case of the TAT-dependent protein in spot 104, we observed that the twin arginines in five orthologs were altered to KR while preserving the other features of a TAT motif. This finding suggested that in some cases TAT substrates also can contain KR. We observed several additional cases showing this phenomenon, such as spot 71. Interestingly, very recently, a similar observation was made with an *E. coli* protein (Hinsley et al., 2001). Further experimentation is needed to confirm that the orthologs with KR in fact translocate via the TAT pathway.

Use of Cross-Correlation Plots as a Systematic Method for the Identification of Post-Translational Modifications, Large Annotation Errors, and Alternative Splicing

Cross-correlation between the theoretical and experimental pI and molecular mass values generally showed very good agreement between theory and experimentation, after removal of the presequences, for both pea and Arabidopsis. However, discrepancies existed for a few spots and suggested dimerization and post-translational modifications. One very clear case of alternative splicing was discovered, and comparison with EST data showed that one of the forms was expressed preferentially in both pea and Arabidopsis. The MALDI-TOF MS data covered the protein sequence nearly completely, thus providing strong evidence that the annotation of the sequence was correct. Interesting alternative splicing changed only the C terminus of the protein, strongly affecting the pI value of this luminal protein. It is unclear if the expression of the acidic (pI 5.3) and basic (pI 9.3) forms relates to the strong fluctuation of pH in the lumen (between 3.5 and 7.0) in conjunction with the day and night cycles. This demonstrates that the cross-correlation plots can help to identify annotation errors that affect pI or molecular mass as well as unexpected splicing events. Moreover, it indicates the presence of positive cleavable presequences as well as oligomerization and certain post-translational modifications.

Evaluation of the Genome-Wide Localization Prediction of the Luminal Proteome

Both TargetP and SignalP have become very popular localization predictors in the plant biology field and are used widely for protein localization and presequence prediction in many published articles (Arabidopsis Genome Initiative, 2000), reviews (Adam et al., 2001), and database entries. A relatively new subcellular localization program is Predotar, which is focused completely on distinguishing between mitochondrial and chloroplast-targeted proteins but makes no cleavage site predictions (<http://www.inra.fr/Internet/Produits/Predotar/>). We observed that the performance of Predotar was significantly less accurate than that of TargetP (Tables 1 to 3), and the program seemed biased toward well-known proteins involved in photosynthesis. SignalP has been used regularly to determine potential thylakoid luminal proteins, although it was developed specifically to identify bacterial and ER signal peptides. In our earlier proteomics study of pea thylakoids, we evaluated the performance of these predictors and speculated that it would be possible to predict a large fraction of the luminal proteome if additional constraints based on experimentally identified luminal proteins and their N termini were included (Peltier et al., 2000; van Wijk, 2000).

In this study, therefore, we extensively explored the use of

these localization predictors (TargetP and SignalP-HMM) in combination with additional constraints derived from (1) a set of 211 experimentally identified known nucleus-encoded chloroplast proteins collected from SWISS-PROT release 38 that are not located in the lumen (this is the test set used to train TargetP; for a complete list, see <http://www.cbs.dtu.dk/services/TargetP/datasets/datasets.html>); (2) 41 experimentally identified nonredundant luminal proteins (from this article and other studies) and 160 orthologs in higher plants and green algae, providing a set of 201 luminal proteins (109 proteins have a typical TAT motif in the ITP); and (3) the use of TMHMM to remove TM proteins.

TargetP predicted 3646 proteins localized in plastids (using all five reliability classes), and 38% have no annotated function in MIPS. On the basis of TargetP prediction of an earlier test set (Emanuelsson et al., 2000), this prediction is expected to be ~70% accurate, with a sensitivity of 85%. Analysis with TM prediction programs indicates that ~500 proteins (45% have no annotation) of these 3646 proteins have at least one TM domain and are located in the thylakoid membrane or in the inner chloroplast envelope. The other 3146 proteins (38% have no annotation) are soluble proteins and are located in the plastid stroma or thylakoid lumen (O. Emanuelsson and G. von Heijne, unpublished data). TargetP correctly predicted chloroplast localization for 89% of the 174 complete luminal orthologs when including the algae and 94% without the green algae. Predotar correctly predicted 67%.

Because there was no reason to select either the Gram-negative or the Gram-positive version of SignalP (based on either performance [Table 1] or theory), we used the union of the predictions for Gram-negative and Gram-positive SignalP-HMM predictions. We progressed stepwise through our prediction strategy while monitoring the accuracy by measuring false positives in the set of 211 known stromal proteins and monitoring the sensitivity by measuring the success rate in the set of 41 known luminal proteins and the set of orthologs (Table 5). The end result was two sets of potential luminal proteins, one targeted through the TAT pathway and the other targeted through the Sec and possibly other pathways. Within each set, proteins were sorted into four groups (p, s, a, and r) based on the amino acids in the -3 and -1 positions directly upstream of the lumen cleavage site. The thylakoid-processing peptidase that is responsible for processing of the ITPs is sensitive to the amino acids in the -3 and -1 positions, as was demonstrated by mutagenesis studies in both higher plants and *C. reinhardtii* (Bassham et al., 1994; Kuras et al., 1995). The prediction process was quite successful, although not perfect, for the proteins with a TAT motif, with a false positive rate of 0.06 and a sensitivity of 0.83 using the permissive filter for the -3 and -1 positions, and 93 luminal proteins with a TAT motif were predicted. Application of more stringent filters reduced the sensitivity and improved the accuracy only slightly. The prediction for the non-TAT proteins was not as strong, with a false positive rate of 0.14 and a

sensitivity of 0.60 using the p motif (Table 5). The better prediction of the TAT proteins is attributable to the presence of the TAT motif, which is a relatively unique signature. When analyzing both sets of predicted proteins manually, the accuracy was lower because of the presence of nonchloroplast proteins in the TargetP prediction. These false positives were not taken into account in the accuracy and sensitivity measurements because these were measured on chloroplast proteins directly.

Given the better performance, we present only the set of predicted TAT proteins. Only 30% of the 93 predicted proteins were annotated with some function in MIPS, and a manual annotation using all major domain predictors increased annotation to 56%. Twenty-two of the 93 predicted proteins were removed manually because functional domains or experimental data (from the literature and the databases) seemed incompatible with their predicted luminal localization. A summary of the functions of the predicted proteins is shown in Figure 10B. Forty-four percent of the proteins have no assigned function, 18% have a role in photosynthesis (some well known and those for the paralogs unknown), 20% function in protein processing, proteolysis, and (un)folding, and 7% function in antioxidative defense and nonphotosynthetic redox reactions. One ABC transporter subunit was predicted, and 10% of the predicted proteins are involved in different metabolic processes.

To improve on the genome-wide prediction for luminal proteins, a specific ITP predictor needs to be trained using the set of 174 luminal proteins. Finally, as discussed elsewhere in this paper, the Arabidopsis gene annotation is erroneous in a significant number of cases, which can affect localization prediction. Thus, careful annotation will help to make a better prediction.

METHODS

Plant Growth

Arabidopsis thaliana, ecotype Columbia, was grown on soil in a temperature-controlled growth chamber under a 10-hr-light/14-hr-dark cycle at 21/16°C light/dark temperatures at a light intensity of $\sim 100 \mu\text{mol}\cdot\text{m}^{-2}\cdot\text{sec}^{-1}$.

Isolation of the Thylakoid Lumen

Leaves from Arabidopsis were homogenized in ice-cold medium A (50 mM Hepes-KOH, pH 7.5, and 100 mM sorbitol supplemented with $50 \mu\text{g}\cdot\text{mL}^{-1}$ Pefablok, $2 \mu\text{g}\cdot\text{mL}^{-1}$ antipain, and $2 \mu\text{g}\cdot\text{mL}^{-1}$ leupeptin), filtered through four layers of Miracloth (Calbiochem; $22 \mu\text{m}$), and centrifuged for 3 min at 9000g. Thylakoid pellets were resuspended in medium A with a cocktail of nine protease inhibitors (antipain [$40 \mu\text{g}\cdot\text{mL}^{-1}$], leupeptin [$5 \mu\text{g}\cdot\text{mL}^{-1}$], pepstatin [$0.7 \mu\text{g}\cdot\text{mL}^{-1}$], Pefablok [$5 \text{ mg}\cdot\text{mL}^{-1}$], E64 [$10 \mu\text{g}\cdot\text{mL}^{-1}$], chymostatin [$20 \mu\text{g}\cdot\text{mL}^{-1}$], bestatin [$40 \mu\text{g}\cdot\text{mL}^{-1}$], phosphoramidon [$5 \mu\text{g}\cdot\text{mL}^{-1}$], and apoprotin [$2 \mu\text{g}\cdot\text{mL}^{-1}$]) and centrifuged on a Percoll gradient (0 to 90%) in me-

dium A for 4 min at 3000g. The thylakoid band was collected and washed three times in the same medium A. Thylakoids were passed twice through a Yeda press (at 100 bar), and thylakoid membranes were removed subsequently by ultracentrifugation (60 min at 150,000g). The membrane-free supernatant was collected and concentrated to $20 \text{ mg protein}\cdot\text{mL}^{-1}$. Alternatively the membrane-free supernatant was concentrated 10-fold and precipitated in 90% acetone and the luminal pellet was resuspended at $20 \text{ mg protein}\cdot\text{mL}^{-1}$.

Construction and Analysis of Denaturing Two-Dimensional Gels

Luminal proteins ($250 \mu\text{g}$ for silver stain and 1 to 1.5 mg for Coomassie Brilliant Blue R 250 stain) were solubilized in 7 M urea, 2 M thiourea, 2 mM Tributyl phosphine, 4% 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonic acid, 0.5% Triton X-100, and 2% pharmalyte, pH 3 to 10 (map pH 4 to 7), or 2% immobilized pH gradient buffer, pH 6 to 11 (map pH 7 to 11) (Rabilloud, 1998). Immobililine DryStrips (Amersham Pharmacia, Uppsala, Sweden) (pH 4 to 7 and pH 6 to 11) were rehydrated overnight in sample suspension at room temperature. Isoelectric focusing in the first dimension, gradient Tricine-SDS-PAGE in the second dimension, and staining and image analysis of the gels were performed as described by Peltier et al. (2000). Expression levels were calculated using the two-dimensional analysis software Melanie (Bio-Rad).

Mass Spectrometry, Database Searching, Sequence Analysis, Localization Prediction, Sequence Alignments, and Matching with Expressed Sequence Tags

Stained protein spots were excised from the gel, washed, reduced, and digested with modified trypsin (Promega) according to Shevchenko et al. (1996). The peptides then were extracted and dissolved in $20 \mu\text{L}$ of 5% formic acid, and $0.5 \mu\text{L}$ was applied to the matrix-assisted laser desorption ionization time-of-flight target plate by the dried droplet method using α -cyano-4-hydroxycinnamic acid as a matrix. When necessary, the samples were concentrated using microcolumns (Gobom et al., 1999) and eluted directly onto the matrix-assisted laser desorption ionization target. The mass spectra were obtained manually or in automated mode using a matrix-assisted laser desorption ionization time-of-flight mass spectrometer (REFLEX II from Bruker Daltonics (Billerica, MA) or Voyager-DE-STR from Applied Biosystems [Foster City, CA]). The spectra were calibrated internally using tryptic peptides from autodigestion and were annotated with the program m/z from Proteometrics (www.proteometrics.com) or automatically using PSI software (Applied Biosystems). The latest versions of the National Center for Biotechnology Information nonredundant database were searched with the resulting peptide mass lists, manually using the search engine ProFound (www.proteometrics.com) or automatically using MS-Fit (Protein Prospector) and PSI solutions (Applied Biosystems). The search strategy was in principle as described previously (Peltier et al., 2000).

To further analyze the samples, the remainder of the extracted peptides were desalted, concentrated on microcolumns (Poros R2 or R3; Applied Biosystems), and eluted directly into nanoelectrospray needles (Protana A/S, Odense, Denmark) with $1.2 \mu\text{L}$ of 50% methanol and 1% formic acid (Wilm et al., 1996). The spectra were acquired on an electrospray tandem mass spectrometer (Q-TOF; Micromass, Manchester, UK). The instrument was calibrated with $1 \mu\text{g}\cdot\text{mL}^{-1}$ Nal in 50% isopropanol. Alternatively, the spectra were

used to search the public databases with the program Mascot (<http://www.matrix-science.com/>). The mass spectrometry spectra usually were interpreted using MassLynx and PepSeq (Micromass), and the resulting sequences were used to search different public databases using FASTA3 (www2.ebi.ac.uk/fasta3).

Manual predictions for chloroplast localization and chloroplast and luminal transit peptides were made using the search engines TargetP (www.cbs.dtu.dk/services/TargetP), SignalP (www.cbs.dtu.dk/services/SignalP), and Predotar (www.inra.fr/Internet/Produits/Predotar).

Data Acquisition and Localization Prediction

Arabidopsis protein sequences for theoretical analyses were obtained on November 30, 2000, via ftp (<ftp://ftpmips.gsf.de/cress/arabiprot/>). The set used in this study is similar but not identical to the *arabi_all_proteins_v211200.tfa* set at this ftp site. This set corresponds to the set presented by the *Arabidopsis* Genome Initiative (2000). Entries At4g341{31, 35, 38} in the set of November 30 have been replaced by At4g34130, and At5g33{390, 400, 410, 420, 430, 440} have changed their names to {W120, C190, C205, C235, W250, W305}ECH54M, respectively.

All 25,460 sequences were processed through subcellular localization predictor TargetP (version 1.01, using plant version; <http://www.cbs.dtu.dk/services/TargetP>) (Emanuelsson et al., 2000), and the 3646 proteins that were predicted to contain a chloroplast transit peptide were processed further through signal peptide predictor SignalP (version 2.0, using HMM version; <http://www.cbs.dtu.dk/services/SignalP-2.0/>) (Nielsen et al., 1997, 1999) and transmembrane predictor TMHMM (version 2.0; <http://www.cbs.dtu.dk/services/>) (Sonnhammer et al., 1998; Krogh et al., 2001). Various constraints were applied (see Results) to obtain the final predicted set of luminal proteins. The scheme of the prediction approach is shown in Figure 10A.

Multialignments and Construction of Phylogenetic Trees

Multialignment and construction of phylogenetic trees were performed using ClustalW (www2.ebi.ac.uk/clustalw/) and MultAlin (<http://www.toulouse.inra.fr/multalin.html>). Phylogenetic trees were calculated using parsimony and neighbor-joining distance methods (Phylip; Department of Genetics, University of Washington, Seattle) with bootstrap values to test the statistical robustness of the trees, essentially as described by Peltier et al. (2001).

Miscellaneous

Protein determinations were performed according to Bradford (1976). Chlorophyll concentrations were determined spectrophotometrically in 80% acetone (Porra et al., 1989).

Accession Numbers

The accession number for the sequence of the moss *Physcomitrella* mentioned in Figure 4 is 7046852; the accession numbers for the soybean ESTs mentioned in Figure 5 are AW508389, BE823905, and BE607927. The SWISS-PROT accession number for the *Arabidopsis* sequence (spot 71) mentioned in Figure 5 is P83050 and is assigned OEC23 related protein.

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NOTE ADDED IN PROOF

During proofreading of this paper, a description of an isomerase with a TAT motif, an isomerase targeted via the Sec (or other) pathway, and other luminal proteins appeared online (Schubert, M., Petersson, U.A., Haas, B.J., Funk, C., Schröder, W.P., and Kieselbach, T. [2002]. Proteome map of the chloroplast lumen of *Arabidopsis thaliana*. *J. Biol. Chem.*, in press).