Plant Molecular Biology 56: 397–412, 2004. © 2004 Kluwer Academic Publishers. Printed in the Netherlands.

Lateral roots affect the proteome of the primary root of maize (Zea mays L.)

Frank Hochholdinger^{1,5,*}, Ling Guo^{2,3} and Patrick S. Schnable^{1,2,3,4}

¹Department of Agronomy, Iowa State University, Ames, IA 50011-3650, USA (*author for correspondence; e-mail frank.hochholdinger@zmbp.uni-tuebingen.de); ²Department of Genetics, Development, and Cell Biology, Iowa State University, Ames, IA 50011-3650, USA; ³Bioinformatics and Computational Biology Graduate Program, Iowa State University, Ames, IA 50011-36506, USA; ⁴Center for Plant Genomics, Iowa State University, Ames, IA 50011-3650, USA; ⁵Present address: Center for Plant Molecular Biology (ZMBP), University of Tuebingen, Auf der Morgenstelle 28, 72076 Tuebingen, Germany

Received 17 August 2004; accepted in revised form 20 September 2004

Key words: lrt1, maize, mutant, proteome interactions, proteomics, roots

Abstract

Lateral roots are initiated from the pericycle cells of other types of roots and remain in contact with these roots throughout their life span. Although this physical contact has the potential to permit the exchange of signals, little is known about the flow of information from the lateral roots to the primary root. To begin to study these interactions the proteome of the primary root system of the maize (*Zea mays* L.) *lrt1* mutant, which does not initiate lateral roots, was compared with the corresponding proteome of wild-type seedlings 9 days after germination. Approximately 150 soluble root proteins were resolved by two-dimensional electrophoresis and analyzed by MALDI-ToF mass spectrometry and database searching. The 96 most abundant proteins from a pH 4–7 gradient were analyzed; 67 proteins representing 47 different Genbank accessions were identified. Interestingly, 10% (15/150) of the detected proteins were preferentially expressed in *lrt1* roots that lack lateral roots. Eight of these *lrt1*-specific proteins were identified and four are involved in lignin metabolism. This study demonstrates for the first time the influence of lateral roots on the proteome of the primary root system. To our knowledge this is the first study to demonstrate an interaction between two plant organs (viz., lateral and primary roots) at the level of the proteome.

Introduction

The root system of maize (*Zea mays* L.) comprises embryonically formed primary and seminal roots, as well as post-embryonic shoot-borne crown and brace roots (Hochholdinger *et al.*, 2004a). A common characteristic of all root types is the formation of lateral roots. Lateral roots play an important role in root stock architecture (Lynch, 1995) and are responsible for the majority of water and nutrient uptake by maize plants (McCully and Canny, 1988; Varney and Canny, 1993; Wang *et al.*, 1991, 1994, 1995). Lateral roots are initiated in the differentiation zone of the main root some distance away from the root apex where pericycle cells are no longer actively dividing (Esau, 1965). Developing main roots constantly proliferate new lateral roots. Thus, the longitudinal positions of lateral roots along the main root indicate their developmental status, with the younger lateral roots located closer to the root tip. The root tip region contains also many newly initiated lateral root primordia that have been initiated but are not yet visible.

Few mutants that affect lateral root formation have been isolated in the monocotyledonous cereals (Hochholdinger *et al.*, 2004a) with only one mutant (*lrt1*) affected in the initiation of lateral roots. The *lrt1* mutant is specifically affected in the early post-embryonic phase in failing to initiate lateral roots from the embryonic primary and seminal roots and crown roots from the coleoptilar node (Hochholdinger and Feix, 1998).

Each individual main root and its lateral roots represents a small 'root module' consisting of hundreds of lateral roots that must interact closely with the main root to which they are connected. Signaling between main roots and lateral roots plays a pivotal role for the formation of the mature root stock (Lynch, 1995). This implies that there must be a considerable flow of information between lateral roots and the main roots. One means by which information is transmitted from the main roots to lateral roots is via polar auxin transport along the main root, which is known to be an essential signal for lateral root initiation (Reed et al., 1998). It can be hypothesized that, lateral roots which make up the majority of the absorbing surface of roots and are therefore instrumental in the perception of environmental cues, may send signals back to the main roots and thereby influence the status of the main root. Although this flow of information from the lateral root system would be expected to influence gene expression and thus protein accumulation in the main roots, little is known about this communication.

In this study the *lrt1* mutant, which does not initiate lateral roots, was used as a model to study the transfer of information from lateral roots to the primary root system. The accumulation of proteins in 9-day-old primary root systems that include normal lateral roots (wild-type) was compared with the accumulation of proteins in primary root systems that do not include lateral roots (lrt1). This study demonstrates for the first time that the absence of lateral roots has a considerable impact on the composition of the proteome of the primary root. Specifically, 10% (15/150) of detected proteins accumulated in *lrt1* primary roots to levels that were at least three times higher than in wild-type primary roots, providing evidence of communication from the lateral roots to the primary root system.

Material and methods

Plant material

The mutant lrt1 was initially isolated from the F₂-generation of an EMS-mutagenized B73

population (Hochholdinger and Feix, 1998). Heterozygous wild-type plants were selfed over three subsequent generations. Primary roots of lrt1 mutant seedlings for the experiments described here were harvested from segregating families of the F₅-generation, while the primary root system of wild-type seedlings was collected from homozygous wild-type families of the F₅-generation. Each primary root protein extract was obtained from a pool of approximately 40 primary roots. Each pool consisted of 10 roots from each of four different families. Biological replicates were generated from independent root samples from different families. Proteins were isolated from complete primary roots that were cut off 1 mm below the coleorhiza and included the primary root tip. Wild-type and *lrt1* seedlings were grown in paper rolls according to Hetz et al. (1996) at 28 °C in dark.

Maize primary root protein isolation

Total proteins were isolated *via* acetone precipitation from primary root systems of 9-day-old wildtype and *lrt1* seedlings according to Damerval *et al.* (1986) as described in Hochholdinger *et al.* (2004b). Dried protein pellets were resuspended in a solution containing 7 M urea, 2 M thiourea, 4% CHAPS, 2 mM TCIP (Pierce, Rockford, IL), 0.5% Bio-Lytes 3/10 (Biorad, Hercules, CA), 40 mM Tris, and 0.001% orange G dye.

Two-dimensional separation of total primary root proteins and MALDI-TOF mass spectrometry

Isoelectric focusing of protein extracts from maize primary roots was performed with the IPG Phor isoelectric focusing unit (Amersham Pharmacia Biotech, Uppsala, Sweden) using 18 cm immobilized, linear pH 4-7 gradients. Proteins were then separated according to their M_{rs} on 12–18% SDS polyacrylamide gradient gels, stained with a modified colloidal Coomassie blue stain and scanned with a Biorad GS 710 scanner (Biorad, Hercules, CA) as described previously (Hochholdinger et al., 2004b). Images were quantified and normalized with Biorad quantity one and PDQuest software prior to comparison of respective spots on wild-type and lrt1 gels. Preparation of the samples for MALDI-ToF mass spectrometry was performed as previously described (Hochholdinger et al., 2004b). Crystallized proteins were analyzed with a VoyagerDE PRO mass spectrometer (PerSeptive Biosystems, Framingham, MA).

Analysis of spectrometric data

Proteins were identified via peptide mass fingerprints using the MS-fit program of the protein prospector package (Clauser et al., 1999; http:// prospector.ucsf.edu/) and databases NCBInr, pdbEST others, the maize EST contig database ZMtuc (www.maizegdb.org) (all databases as of 03.26.2004) and assembled genomic sequences (MAGIs) from Version 3.1 of the ISU maize genome assembly project (www.plantgenomics.iastate.edu/maize). Mass spectrometry utilities (MSU) software was used to automate the MS-fit identification tools (Porubleva et al., 2001). A protein identified via the MS-fit database search was accepted as correct only if all criteria defined by Hochholdinger et al. (2004b) were met (see footnotes of Table 1). Sequence similarity searches for EST and genomic sequences were performed using the blastx program (Altschul et al., 1997).

Comparisons of different maize proteomes

Maize protein sequences identified from the NCBInr protein database were directly compared to the maize proteins from the datasets of Chang *et al.* (2000) and Porubleva *et al.* (2001) as described in Hochholdinger *et al.* (2004b). The protein sequences from Chang *et al.* (2000) and Porubleva *et al.* (2000) and Porubleva *et al.* (2001) were downloaded from Genbank using the protein accessions provided in these manuscripts. The maize protein dataset from the current study was set as the query file to search these two maize data sets using blastp with an *E*-value cut-off of 1e-10.

Results

purification and solubilization of maize primary root proteins

Total soluble root proteins were isolated from the primary root systems of 9-day-old wild-type and *lrt1* maize seedlings. At this stage, lateral roots had already formed on the wild-type primary roots (Figure 1). Consistent with the known phenotype (Hochholdinger and Feix, 1998), seedlings homo-

zygous for the lateral root initiation mutant *lrt1* did not form any lateral roots. Isoelectric focusing of total soluble primary root protein extracts was performed on a linear gradient: pH 4-7. After isolelectric focusing, proteins were separated according to their masses in a second dimension and stained with Coomassie blue. A total of 150 protein spots were visible including several very faint spots, which were not analyzed spectrometrically. Each map was made in triplicate from independent primary root protein preparations and excellent reproducibility was obtained (data not shown). The comparative two-dimensional electrophoresis maps of wild-type and *lrt1* primary root proteins are shown in Figure 2. These maps are synthetic match sets created with PDQuest software. Each map is composed of the three gels for wild-type and *lrt1* protein extracts and represents average spot intensities as calculated by PDQuest after filtering and smoothening the original scans. The intensities of all spots on a gel were normalized in silico with PDQuest to compensate for non-expression related variations in spot intensity. Average normalized spot intensities from the gels are provided in Table 1. Overall, excellent reproducibility of spot intensities was obtained among the three replicates of a genotype; the average coefficient of variance per spot for wild-type and *lrt1* gels was 42% and 35%, respectively, which are similar to those observed by Chang et al. (2000) and in a previous dataset from our laboratory comparing maize mitochondrial N- and T-cytoplasm proteins (38% and 39%) (Hochholdinger et al., 2004b). As expected, less abundant proteins often had higher % CVs.

Identification and functional annotation of primary root proteins

The 96 most abundant proteins were eluted from a representative two-dimensional gel of the mutant *lrt1*, digested with trypsin and analyzed using a MALDI-ToF mass spectrometer according to the strategy described in Hochholdinger *et al.* (2004b). Automated MS-fit software was used for searching protein databases on all available higher plant proteins (streptophyta) since the maize genome has not yet been completely sequenced and many genes are highly conserved among higher plants.

Sixty seven of the 96 analyzed proteins were identified by matching known proteins or translated

Table 1. Primary root against the NCBInr pro	proteins fro otein db, the	m wild-typ e ZMtuc m	e and <i>lrt1</i> 1 aize EST c	naize seedlings ontig db, the E	identified after two ST singletons of th	o-dimensional separatior re dbEST.others db, and	and MALDI-ToF analys the ISU maize genomic set	sis of trypsi equences a	in-digested tssembly dl	b (MAG	s matched I) entries.
Spot No.	MOWSE score ^a	Peptides matched (n) ^b	% EST/ protein covered ^e	$M_{\rm r}$ predicted/ $M_{\rm r}$ gel ^d	p <i>I</i> predicted/p <i>I</i> gel ^e	EST AC [species] library [/]	Function (blastx hit) [snecies] AC ^g	Intensity ^h	i.		
		Ì						ΜT	% C.V.	lrt]	% C.V.
Cell growth/division ⁱ											
60	2.17e+4	Ś	44	40.3/49.2	4.59/4.56	BE575292 [Z. mays] tassel primordium	RAD23 protein [D. carota] Y12014	3043	50	5875	21
Cellular organization											
15	2.04e+5	9	24	37.1/45.7	5.46/5.31	ZMtuc03-09-11.11965	Actin [Z. mays] AAB40103	27060	29	27461	33
16	8.35e + 5	11	39	37.1/45.6	5.46/5.37	ZMtuc03-09-11.11965	Actin [Z. mays] AAB40103	57408	50	25111	24
17	4.72e+5	٢	27	37.1/45.6	5.46/5.37	ZMtuc03-09-11.11965	Actin [Z. mays] AAB40103	15755	6	14675	18
Disease/defense											
32	1.34e + 4	4	34	13.1/16.5	4.55/4.46	AW600568 [Z. mays]	Copper chaperone	5707	86	8556	24
						mixed adult tissues	homolog CCH				
							[O. sativa] T50779				
41	8.18e + 4	6	43	27.2/25.9	5.42/6.26	CD438839 [Z. mays]	L-ascorbate peroxidase	44999	27	32437	48
						endosperm	[O. sativa] 103595				
42	1.49e + 6	9	40	27.2/26.1	5.42/5.72	CD438839 [Z. mays]	L-ascorbate peroxidase	40806	б	33041	32
72	7 849 + 4	v	90	0969 26	5 10/5 70		U. Sunval 100.001 [U. Sundaced	5405	02	74384	19
2	2	<i>,</i>	2	0.01 0.11		2 mm ear	[H. vulgare] AAL08496	2	2	-	
44	3.56e + 4	5	21	27.2/26.0	5.42/5.91	CD438839 [Z. mays]	L-ascorbate peroxidase	12623	50	21887	31
						endosperm	[O. sativa] T03595				
56	1.80e + 3	5	18	32.9/36.5	5.70/5.80	NCBI	Sulfur starvation	8859	6	9133	23
							induced [Z. mays] U33318				
70	3.41e+3	5	25	25.5/32.9	7.11/5.04	NCBI	Superoxide dismutase-3	23792	59	12337	55
							precursor [Z. mays] X12540				
76	3.45e + 3	4	18	27.3/26.0	5.28/5.43	ZMtuc03-08-11.13269	L-ascorbate peroxidase [Z. mays] S49914	11187	94	4298	14

Spot No.	MOWSE score ^a	Peptides matched	% EST/ protein	$M_{\rm r}$ predicted/ $M_{\rm r}$ gel ^d	p <i>I</i> predicted/p <i>I</i> gel ^e	EST AC [species] library ⁶	Function (blastx hit) fsneciesl AC ⁶	Intensity ^h			
		(i)						WT	% C.V.	lrtl	% C.V.
Energy 10	1.69e+6	10	37	59.1/50.9	6.01/5.11	NCBI	F-1-ATPase subunit	25082	5	3950	71
11	9.42e+6	11	37	59.1/50.8	6.01/5.20	NCBI	2 [Z. mays] M36087 F-1-ATPase subunit	59850	4	26736	12
				-	-		2 [Z. mays] M36087				
12	8.68e+8	13	32	59.1/50.6	6.01/5.29	NCBI	F-1-ATPase subunit	116636	29	47793	27
13	9.47e+7	13	35	59.1/50.6	6.01/5.36	NCBI	F-1-ATPase subunit	13014	59	20249	21
							2 [Z. mays] M36087				
14	3.85e+6	10	26	59.1/50.6	6.01/5.25	NCBI	F-1-ATPase subunit	25813	18	29220	17
0		4	5				2 [Z. mays] M36087		10	0.4 50	č
49	1.3/e + 6	10	51	30.4/38.9	1.02/0.72	NCBI	Cytosolic ølvceraldehvde-3-	1630/	48	9458	94
							phosphate				
							dehydrogenase				
							GAPC3 [Z. mays]				
							U45856				
50	4.35e + 4	8	24	36.4/40.3	7.02/6.51	NCBI	Cytosolic	12725	25	19876	23
							glyceraldehyde-3-				
							phosphate				
							dehydrogenase				
							[Z. mays] U45856				
51	1.73e + 5	8	29	36.5/40.6	6.40/6.57	NCBI	Glyceraldehyde-3-	9838	67	11398	48
							phosphate				
							dehydrogenase				
							[Z. mays] U45858				
52	1.91e + 6	6	39	36.5/41.7	6.40/6.46	NCBI	Glyceraldehyde-	20566	22	17789	18
							3-phosphate				
							dehydrogenase				
							[Z. mays] U45858				
65	2.04e+3	4	19	19.7/18.6	5.19/4.93	ZMtuc03-08-11.5252	Mitochondrial F0	4125	64	797	22
							ATP synthase D chain [0. sativa] BAD09007				

Table 1. Continued.

Table 1. Con	tinued.				
Spot No.	MOWSE	Peptides	% EST/	$M_{ m r}$ predicted/	$\mathrm{p}I$
	score ^a	matched	protein	$M_{ m r}~{ m gel}^d$	pred
		$q(\mathbf{u})$	$covered^c$		

Spot No.	MOWSE score ^a	Peptides matched (n) ^b	% EST/ protein covered ^c	$M_{ m r}$ predicted/ $M_{ m r}$ gel ^d	p <i>I</i> predicted/p <i>I</i> gel ^e	EST AC [species] library [/]	Function (blastx hit) [species] AC ^g	Intensity ^h			
		Û						ΜT	% C.V.	lrtI	% C.V.
99	4.74e+7	12	48	19.7/18.4	5.19/5.11	ZMtuc03-08-11.5252	Mitochondrial F0 ATP synthase D chain [0. sativa] BAD09007	3343	41	3041	34
Metabolism											
18	6.30e+3	5	25	39.2/44.2	5.34/5.33	ZMtuc03-08-11.28030	Glutamine synthetase [Z. mays] BAA03430	12800	64	14675	18
19	3.43e+3	9	14	39.0/44.6	5.23/5.30	NCBI	Glutamine synthetase [Z. Mays] X65929	15457	8	9213	13
39	1.64e + 3	4	22	16.6/15.9	6.30/6.47	ZMtuc03-08-11.11114	Nucleoside diphosphate	3124	36	7511	24
							kinase [<i>S. officinarum</i>] P93554				
53	2.71e+5	16	26	58.4/54.2	5.52/6.50	NCBI	Beta-Glucosidase	11592	20	22018	21
54	6.01e+6	15	23	58.4/54.4	5.52/5.68	NCBI	Beta-Glucosidase	8763	38	22417	62
				-			[Z. mays] 1E4LA				
58	1.79e+6	8	32	27.4/28.8	5.46/4.59	ZMtuc03-08-11.6743	Methylthioadenosine/ S-adenosyl	6085	32	8088	11
							homocysteine nucleosidase [O. sativa]				
ç			9				AAL58883	000	c t		Į
60	2.99e + 3	4	18	21.1/21.9	0.00/0.02	21VILUCU3-U3-U3-11.82U9	Futative	665	<i>c</i> /	4400	10
							1,4-061204untore reductase [<i>O. sativa</i>] BAB92583.1				
71	6.73e + 4	5	30	21.7/21.5	6.06/6.39	ZMtuc03-08-11.8209	Putative	2251	36	4367	62
							1,4-benzoquinone reductase [0. sativa] BAB92583.1				
72	4.72e+3	Ś	23	23.9/23.5	5.96/6.49	NCBI	Glutathione transferase III(b) [Z. mays]	10256	74	3592	41
							AJ010296				
82	4.39e+4	8	22	33.4/35.0	5.72/4.99	ZMtuc03-08-11.29884	Putative inorganic	8669	32	5131	12
							pyrophosphatase [A. thaliana] NP_196527				

Table 1. Continued										
Spot No.	MOWSE score ^a	Peptides % matched pr	6 EST/ totein	$M_{ m r} { m predicted}/{M_{ m r}} { m gel}^d$	p <i>I</i> predicted/p <i>I</i> gel ^e	EST AC [species] [ibrary [/] (Function (blastx hit) [suecies] A C ^g	Intensity ^h		
			parate			_		WT	% C.V.	IrtI % C.V.
87	5.38e+4	8 14	4	63.1/58.9	5.46/5.68	NCBI	Phosphoglucomutase 1 [Z. mays] U89341	3261	39	12689 20
88	2.46e+3	7 12	0	60.6/57.1	5.29/5.57	NCBI	2,3-bispho- sphoglycerate-in- denendent nhoenhoody	6468	17	7781 36
						. ,]	cerate mutase [Z. mays] M80912			
89	3.40e+3	8 13	ŝ	60.6/56.9	5.29/5.69	NCBI	2,3-bispho- sphoglycerate-in- dependent	12686	61	10598 11
							phosphoglycerate mu- tase [Z. mays] M80012			
92	7.32e+5	15 23	~	58.4/54.6	5.52/5.58	NCBI	Beta-Glucosidase	3491	69	13780 34
93	2.81e+3	6 32	6	39.3/43.5	5.60/5.60	ZMtuc03-08-11.5416	[Z. muys] 1E4LA Glutamine synthetase	10011	18	3876 4
96	1.08e + 3	5 15		32.4/35.4	5.59/5.95	ZMtuc03-08-11.10346	[Z. mays] P38559 Glvoxalase I [Z. mavs]	2129	20	5013 87
						1	AAP76396			
Protein destination and folding	2									
7	2.22e+4	9 20	0	57.1/57.1	5.24/4.78	NCBI	Protein disulfide isomerase [Z. mays] 1.39014	6446	26	20087 24
ი	5.74e+4	9 18	~	57.1/57.2	5.24/4.87	NCBI	Protein disulfide isomerase [Z. mays] 139014	6649	25	12295 20
Ŷ	4.82e+3	5 22	2	57.4/55.0	4.38/4.94	AV915993 [<i>H. vulgare</i>] shoots] 1 5	RuBisCO subunit binding-protein alpha subunit [<i>T. aestivum</i>]	9150	34	4395 31

Spot No.	MOWSE score ^a	Peptides % matched p	6 EST/ <i>I</i> rotein <i>A</i>	$M_{\rm r}$ predicted/ $M_{\rm r}$ gel ^d	/ pI predicted/pI gel ^e	EST AC [species] library [/]	Function (blastx hit)	Intensity [/]	~	
		c (II)	overed				[species] AC	ΤW	% C.V	lrt1 % C.V.
Protein synthesis 31	3.06e+4	4 9	5 1	7.4/19.8	10.43/4.52	BF480867 [<i>S. propinquum</i>] floral meristems	40S ribosomal protein S15 [E. oleifera] AE404770	1962	54	8747 65
Secondary metabolisn 4	n 9.68e+3	4	5	5 9. 8/54.8	9.20/4.84	BI955765 [H. vulgare] seedling	Cinnamate Cinnamate 4-hydroxylase CYP73 [C. sinensis]	3834	59	2647 22
28	7.96e+3	4	3	1.9/32.8	5.10/4.89	ZMtuc02-12-23.6818	AAF66066.2 Caffeoyl-CoA 3-O-methyltransferase [0. sativa] BAA78733	7954	62	35445 34
Transcription 36	4.04e+4	3	6 1	5.5/14.9	8.01/4.75	CD970259 [Z. mays] mixed adult tissues	Glycine-rich RNA-binding protein	6132	118	18240 8
38	8.08e+5	7 5	9 1	3.7/16.0	9.34/5.21	CD661992 [Z. <i>mays</i>] 2 mm ear	2 [0. satival NP_914833 Glycine-rich RNA-binding protein 1	5284	67	27933 38
94	3.52e+3	4	3	8.8/41.4	4.85/5.58	MAGI_43849	 [Z. mays] Q99069 Transcription factor EREBP-like protein [C. arietinum] CAD56217 	2313	65	5308 15
Transport 7	1.25e+6	11 2	5 0	;4.1/52.6	5.07/5.04	NCBI	Vacuolar ATPase B subunit [0. sativa]	20504	11	3814 49
∞	3.05e+10	0 15 2	e v	: 4.1/52.6	5.07/5.11	ZMtuc03-08-11.6793	AB055106 Vacuolar ATPase B subunit [<i>O. sativa</i>] AAK54617	19958	21	14973 9
6	5.78e+9	14 2	9 9	54.1/52.4	5.07/5.18	ZMtuc03-08-11.6793	Vacuolar ATPase B subunit [<i>O. sativa</i>] AAK54617	15314	11	16234 20

Table 1. Continued.

Table 1. Con	tinued.										
Spot No.	MOWSE score ^a	Peptides matched	% EST/ protein	$M_{ m r}$ predicted/ $M_{ m r}$ gel ^d	p <i>I</i> predicted/p <i>I</i> gel ^e	EST AC [species] library	Function (blastx hit) [snecies] AC ^g	Intensity ^h			
		Ì						WT	% C.V.	lr t l	% C.V.
73	1.13e + 5	9	57	26.8/24.2	8.86/5.16	BG840471 [Z. mays] seedling and silk	Chloroplast Cpn21 protein [<i>A. thaliana</i>] NM 122079	4080	73	8713	37
86 Unclassified	3.91e+5	10	18	62.0/59.1	5.89/5.57	NCBI	Vacuolar ATPase 69 kDa subunit [Z. <i>mays</i>] U36436	4618	59	10620	28
21	1.05e + 9	11	39	34.8/41.5	4.90/4.85	ZMtuc03-08-11.5167	Similar to late embryogenesis abundant proteins [O. sativa] AAS07355	5055	40	11059	32
22	2.75e+7	6	32	34.8/41.2	4.90/4.92	ZMtuc03-08-11.5167	Similar to late embryogenesis abundant proteins [O. sativa] AAS07355	5453	67	4394	41
29	7.80e+3	2	31	18.7/21.8	4.52/4.38	ZMtuc03-08-11.15619	Translationally controlled tumor protein homolog [7, mays] AAN40686	5091	16	7163	17
30	7.81e+3	S	31	18.7/21.8	4.52/4.45	ZMtuc03-08-11.15619	Translationally controlled tumor protein homolog [Z. mays] AAN40686	3071	12	2576	63
46	7.19e + 5	7	28	41.7/38.8	6.25/6.49	MAGI 79568	Putative r40c1 protein [0. sativa] NP 912421	2439	77	8096	50
47	7.14e+5	×	34	41.7/38.8	6.25/5.59	MAGI 79568	Putative r40c1 protein [O. sativa] NP_912421	4646	70	0866	7
Unknown 26	1.75e + 7	10	40	n.p./37.7	n.p./4.81	BE129913 [Z. mays] mixes adult tissue	None	15650	30	17161	51
27	3.98e+4	4	91	n.p./33.3	n.p./4.85	MAGI_19154	None	38865	47	18019	18

Table 1. Continued.

Continued.	
Ι.	
Table	

Peptides % EST/ $M_{\rm r}$ predicted/ matched protein $M_{\rm r}$ gel ^d $(n)^b$ converse.	% EST/ $M_{\rm r}$ predicted/ protein $M_{\rm r}$ gel ^d	$M_{ m r}$ predicted/ $M_{ m r}$ gel ^d		p <i>I</i> predicted/p <i>I</i> gel ^e	EST AC [species] library ^f	Function (blastx hit) [suecies] A C ^g	Intensity ^h			
	(11)					Ar formade	WT	% C.V.	lr t l	% C.V.
3+3	4	51	n.p./15.9	n.p./5.35	BI396264 [Z. mays] juvenile shoots	None	4145	4	7366	0
6+3 6	4	43	n.p./15.8	n.p./5.49	BI431138 [Z. mays] juvenile shoots	None	1724	29	2470	72
e + 4	4	56	n.p./18.8	n.p./6.05	B180125 [O. sativa] endosperm	None	25611	30	15753	34
e + 3	4	68	n.p./27.2	n.p./5.29	AI629731 [Z. mays] leaf primordium	None	4825	26	2391	78
e+3	5	14	20.7/25.8	8.81/4.71	ZMtuc03-08-11.5820	Unknown protein [O. sativa] BAB63543	2079	21	2504	99
e + 4	9	51	n.p./37.0	n.p./4.89	ZMtuc03-08-11.5304	None	6743	62	2665	71

^aMOWSE score: statistical probability of true positive identification of predicted proteins (cut off value: 1e + 3) calculated *via* MS-fit software. Maximum allowed M_r deviation of experimental and predicted peptide fragments : 50 ppm. Allowed missed cleavage: 1.

^bPeptides matched (*n*): number of peptides matching predicted protein sequences (cut-off value: n = 4). ^{c%} EST/protein covered: percentage of predicted protein sequence covered by matched peptides (cut-off value: 10%). ^dM_r predicted/M_r gel: molecular mass of predicted protein on gel (M_r gel = M_r predicted \pm 20%); n.p.: no prediction (proteins without functional annotation).

^epl predicted/pl gel: isoelectric point of predicted proteins/of proteins on gel. ¹EST, EST contig (ZMtuc) or genomic sequence (MAGI) accessions of predicted protein; NCBI: proteins identified *via* the annotated NCBInr protein database. ^gFunction: for ESTs obtained via blasts; for proteins predicted via NCBI according to functional NCBI annotation (cut-off value: 1e-10).

^bbpot intensity: spot intensities of wild-type and *lrt1* proteins after normalization; %C.V.: coefficient of variance = (s/x) * 100; x: standard deviation; x: arithmetric mean value. Functional category: classification of the proteins according to the *Arabidopsis* MATDB (http://mips.gsf.de/proj/thal/db/; Schoof *et al.*, 2002).



Figure 1. Root system of 9-day-old wild-type and *lrt1* seedlings (a) left: wild-type, right: *lrt1*. Close up (b) showing lateral root formation in the wild-type seedling (left) and absence of lateral roots in the *lrt1* mutant (right).

ESTs from plants *via* a combination of mass spectrometry and database searches representing 47 different Genbank accessions. Table 1 summarizes the results of the MS fit database searches. The proteins are sorted according to their functions. Apparent molecular masses (M_r) and isoelectric points (pI) of the proteins according to the position of the proteins on the gel (Figure 2) are compared with theoretically calculated molecular masses. The function of the EST and genomic sequences was evaluated by blastx database searches. It was possible to identify 43 of the 47 accessions by known maize proteins or maize ESTs. The remaining proteins were identified *via* sequence similarity to proteins or ESTs from other monocotyledonous plants.

For most proteins identified *via* the annotated NCBInr database (Table 1) a function could be immediately predicted. No function could be proposed, however, for most proteins identified *via* ESTs (Table 1). These ESTs of unknown function were subjected to blastx searches to assign a function to them. Overall, functions could be assigned to 39 of the 47 identified Genbank accessions.

Proteins preferentially accumulated in lrt1 primary roots

The primary purpose of this study was to identify proteins that preferentially accumulated in

primary roots that lack lateral roots (*lrt1*). Proteins that exhibit such preferential accumulation in this experiment are presumably responding to direct or indirect signals from the lateral roots in wild-type plants. Protein accumulation of identified spots on normalized gels was quantified and the average intensity of these spots on three replicate gels was compared between wild-type and *lrt1* primary roots.

The wild-type primary roots used in these analyses included lateral roots (i.e., lateral roots were not manually excised prior to protein extraction). The presence of lateral root proteins in the wild-type samples could potentially cause two problems. First, proteins that accumulate to very high levels in lateral roots could potentially be identified as being differentially expressed in wildtype vs. lrt1 mutant proteomes. This was not a serious concern because our goal was instead to identify proteins that accumulated preferentially in the absence of lateral roots (i.e., that were upregulated in the *lrt1* mutant). The second potential problem with the presence of lateral root proteins in the wild-type root samples is that of 'dilution'. Specifically, if the lateral roots comprise a large proportion of the root system, lateral root proteins could reduce the relative contribution of some primary root proteins in the wild-type samples to the degree that these proteins would appear to be



Figure 2. Comparative proteomic two-dimensional maps of CHAPS-soluble proteins extracted from 9-day-old maize primary roots of wild-type and the lateral root initiation mutant *lrt1* seedlings showing average spot intensities. Maps for each genotype were prepared from three two-dimensional gels conducted using three independent protein preparations. Proteins were separated in the first dimension according to their pIs on IPG strips pH 4–7 and in the second dimension according to their molecular masses on linear 12–18% SDS-polyacrylamide gradient gels. Proteins were stained with colloidal Coomassie blue G250. Protein spots that were identified *via* MALDI-ToF spectrometry are numbered on the maps. Proteins preferentially accumulated in *lrt1* primary roots are numbered on the *lrt1* gel. Normalized data from these maps are summarized in Tables 1 and 2.

up-regulated in the *lrt1* samples. To address this concern, we determined the contribution of lateral roots to the wild-type root samples. The primary

and lateral roots of wild-type seedlings had average fresh weights of 104 mg (\pm 19 mg) and 9 mg (\pm 6 mg) per seedling, respectively. Hence, the

lateral root proteins would be expected to contribute on average only to $\sim 8\%$ of the wild-type root proteome. Hence, the 'dilution effect' described above would be expected to have only a modest impact on the relative abundance of proteins from the wild-type and *lrt1* samples. We dealt with the potential 'dilution effect' of the wild-type root samples with lateral root proteins by considering proteins as accumulating preferentially only if their abundance levels in the *lrt1* proteome were at least three times higher than in the wild-type proteome and if they did not have overlapping standard deviations.

Even using this conservative 3-fold cut-off level, 15 of the 150 detected proteins accumulated preferentially in *lrt1* primary roots (Table 2). Hence, 10% (15/150) of the detected proteins accumulated preferentially in the primary roots of *lrt1* mutant seedlings that do not accumulate lateral roots.

Eight of these 15 proteins could be identified. They include metabolic proteins (1,4-benzochinone reductase, caffeoyl-CoA-3-*O*-methyltransferase, beta-glucosidase, phosphoglucomutase1) as well as proteins potentially involved in transcription (glycine-rich protein), related to plant defense (Lascorbate peroxidase), and protein synthesis and destination (40S ribosomal protein S15; protein disulfide isomerase). Four of the eight proteins that accumulate preferentially in the *lrt1* primary roots may be related to lignin metabolism (caffeoyl-CoA-3-*O*-methyltransferase, beta-glucosidase, L-ascorbate peroxidase, 1,4-benzochinone reductase).

Three proteins (different F- and V-type ATPase subunits: spots 7, 10, 65) accumulated preferen-

tially in wild-type primary roots. Because the primary roots of wild-type seedlings contained lateral roots these proteins might simply be preferentially expressed to high levels in lateral roots. For this reason and because the analysis of such proteins was outside the scope of this study, they will not be discussed further.

Comparison of the maize primary root proteome with other datasets from maize

The sequences of the maize primary root proteins presented in Table 1 were compared to the sequences of the only two other proteome data sets that are available for maize seedlings (Table 3). Five of 33 distinct proteins (10 protein spots) from the Chang et al. (2000) root tip data set exhibited a high degree of similarity to one or more proteins in our total primary root data set. Similarly, among 109 distinct proteins 20 (represented by 45 spots) from the Porubleva et al. (2001) green seedling leaf data set exhibited a high degree of similarity to one or more proteins in our total primary root data set. Hence, 15% of the root tip proteins identified by Chang et al. (2000) and 18% of the leaf proteins identified by Porubleva et al. (2001) were similar to proteins detected in the 9 dag maize primary root proteome. Homologs of five of 47 proteins identified in our primary root extract were found in both, the proteomes of root tips and green leaves of maize seedlings. Hence, in this maize primary root proteome dataset, 57% of the proteins (i.e., 27 proteins) encoded by distinct Genbank accessions

Table 2. Proteins that preferentially accumulated in *lrt1* primary roots^a.

Spot No. ^b	Specificity ^c	Function (AC)
69	16.4	Putative 1,4-benzoquinone reductase BAB92583.1
38	5.3	Glycine-rich RNA binding protein Q99069
43	4.5	L-ascorbate peroxidase AAL08496
31	4.5	40S ribosomal protein S15 AF404770
28	4.5	Caffeoyl-CoA 3-O-methyltransferase BAA78733
92	4.0	Beta-Glucosidase 1E4LA
87	3.9	Phosphoglucomutase1 U89341
2	3.1	Protein disulfide isomerase L39014

^aProteins were only considered to be differentially expressed if they accumulated to levels that were at least 3-fold different and their standard deviations did not overlap. Three proteins accumulated preferentially in the wild-type primary root systems. All of these proteins were identified as different F- and V-type ATPase subunits (spots 7, 10, 65).

^bThe unidentified spots 1, 34, 55, 61, 63, 90 and 91 also accumulated preferentially in *lrt1* roots.

^c Specificity indicates the ratio of accumulation of a particular protein between *lrt1* vs. wild-type primary root protein preparations.

This study	Function	Maize root tips Chang et al., (2000)	Maize leaves Porubleva <i>et al.</i> , (2001)
10, 11, 12, 13, 14	F-1-ATPase subunit 2	11	1, 2, 26, 27, 28, 29, 30, 31, 201
15, 16, 17	Actin	4	37, 38, 39, 40, 56, 57, 58 ^a
49, 50, 51, 52 ^a	Glyceroldehyde-3-phosphate dehydrogenase	5, 6, 7, 13 ^a	206, 207, 208, 209, 210, 211 ^a
88, 89	2,3-bisphosphoglycerate-independent phosphoglycerate	14, 20, 27, 38	7
6	RuBisCO subunit binding-protein alpha subunit		14
7	Vacuolar ATPase B subunit		23
18, 19, 93 ^a	Glutamine synthetase		43
26, 81	Unknown		70, 71, 72, 143
32	Copper chaperone homolog CCH		160
36, 38 ^a	Glycine-rich RNA-binding protein		73, 74, 75, 77, 90, 165, 168, 169, 170
39	Nucleoside diphosphate kinase		229
41, 42, 43, 44, 76 ^a	L-ascorbate peroxidase		102, 103
73	Chloroplast Cpn21 protein		123, 127

Table 3. Similar proteins identified in the proteomes of maize seedlings.

^a These proteins represent more than one Genbank accession.

were not detected in either of the two other maize seedling proteomes.

Discussion

Analysis of the root proteome of maize

To date, the analysis of differential gene expression in maize roots has mainly been restricted to the analysis of a few genes identified via various methods of differential gene expression on the RNA level (e.g. Goddemeier et al., 1998; Matsuyama et al., 1999a, b; Ponce et al., 2000; Bruce et al., 2001). In this study, the soluble proteomes of 9-day-old maize primary roots harvested from wild-type and *lrt1* seedlings which are defective in lateral root initiation were analyzed. Sixty-seven proteins, which represent 47 different Genbank accession numbers, were identified. The minimal overlap (only five shared proteins) between the major proteins identified in the primary root tip dataset of Chang et al., (2000) and our whole-root dataset might be due to the influence of the differentiation status on protein accumulation; while the root tip contains mainly meristematic tissues, the 9-day-old primary root consists predominantly of differentiated tissue. Given the relative small number of proteins identified in these studies, however, this difference might also be due to the different solubilization techniques used in the two studies.

Absence of lateral roots in ltt1 modulates protein accumulation in the primary root

The architecture of the primary root of wild-type and *lrt1* plants differ in that the *lrt1* mutant does not initiate any lateral roots on the primary root. This morphological difference is also reflected in the protein accumulation profiles of wild-type and mutant primary roots. 10% of the proteins detected in our study (15/150) preferentially accumulate in *lrt1* roots.

The increased abundance of these proteins in the primary root of a mutant that does not initiate lateral roots indicates that direct or indirect communication between the lateral roots and the primary root regulates the proteome of the primary root.

The finding that several abundant proteins are significantly up-regulated in the mutant *lrt1* could be explained by the following model: signals originating in the lateral roots of wild-type seed-lings might either directly or indirectly repress certain genes in the primary root. According to this hypothesis, the signaling pathway between lateral and primary roots is interrupted in *lrt1* due to this mutant's lack of lateral roots and that usually suppress the transcription of certain genes in the primary root are missing in *lrt1*. In the absence of repression, these genes are transcribed and translated in the primary root of *lrt1*.

The casparian strip of the endodermis, which provides a barrier to the apoplastic transport of water and solutes (Hose et al., 2001), is composed of primary cell wall components including lignin. Plants are thought to alter the chemical compositions of their casparian strip during development or in reaction to changing environmental conditions, thereby fine-tuning its resistance to the radial flow of water and nutrients (Hose et al., 2001). Hence, given the role of the lateral roots in the uptake of water and nutrients, the casparian strip is a potential target for the signals that originate in the lateral roots and that modulate the proteome of the primary root (Hose et al., 2001). These signals could be involved in the modification of the chemical composition of the casparian strip (e.g., the lignins) to alter its capacity for the uptake of water and nutrient.

Consistent with this hypothesis, four of the eight identified proteins that were shown to accumulate preferentially in the *lrt1* mutant can be linked to lignin metabolism. A caffeoyl-CoA 3-O-methyltransferase accumulated to levels 4.5 times higher in the *lrt1* primary root system than in the wild-type primary root system. This enzyme plays a crucial role during the syntheseis of monolignols which later polymerize to form lignin (Boerjan et al., 2003). A beta-glucosidase accumulated 4.0 times higher in the lrt1 primary roots. After synthesis of the monolignols these lignin precursors are transferred into an inactive storage or transport form monolignol-4-O-beta-D-glucosides. designated Beta-glucosidases regulate storage and mobilization of these monomers for lignin biosynthesis (Boerjan et al., 2003). An L-ascorbate peroxidase preferentially accumulated in *lrt1* primary roots. After transport and activation of the lignin monomers to the cell wall, lignin is formed through dehydrogenative polymerization of the monolignols (Boerjan et al., 2003). Peroxidases are one class of enzymes thought to be involved in this process (Boerjan et al., 2003). Plants contain multiple peroxidases, e.g., the Arabidopsis genome contains 73 different peroxidase genes (Boerjan et al., 2003); it is not yet clear, however, which are involved in dehydrogenative polymerization. In wood-rotting fungi, peroxidases are involved not in lignin biosynthesis, but in the degradation of lignin (Akileswaran et al., 1999). Although it is not known whether endogenous plant peroxidases are also involved in the degradation of lignin, it appears likely that peroxidases are involved in several aspects lignin metabolism in plants. Finally, a putative 1,4benzoquinone reductase was expressed 16.4 times higher in *lrt1* roots as compared to wild-type roots. Interestingly, this enzyme is known to play a pivotal role in lignin degradation by wood-rotting fungi *via* quinone intermediates (Akileswaran *et al.*, 1999). Significantly, many of the enzymes involved in lignin biosynthesis and degradation have multiple isoforms that are differentially regulated during development (Boerjan *et al.*, 2003).

This is the first study to demonstrate that the absence of lateral roots has an impact on the accumulation of proteins in primary roots. Further analyses of the differentially accumulated proteins identified in this study might reveal insights into the regulation of the molecular processes involved in the interactions between lateral and primary roots. Such analyses might also help to define the developmental regulation of lignin metabolism and the molecular regulation of the composition and behavior of the casparian strip.

Acknowledgements

We thank Kent Vander Velden for advice on using his 'mass spectrometry utilities' program, Ryan Brenke (ISU) and Dieter Steinmetz (University of Tuebingen) for bioinformatic support and Michael Schwall, (Aventis Crop Science), Guenter Feix, (University of Freiburg) and Joseph Dubrovsky (University of Cuernavaca) for helpful discussions. F.H.'s research at Iowa State University was supported in part by a post-doctoral fellowship from the DAAD (German Academic Exchange Service) and by the Suedwestdeutsche Saatzucht, Rastatt, Germany. Additional support was provided by Hatch Act and State of Iowa funds. Current root research in F.H.'s laboratory is supported by the German Scientific Foundation (DFG; award HO2249/4-1), the SFB 446 and the framework program 'heterosis in plants' (award HO2249/6-1).

References

Akileswaran, L., Brock, B.J., Cereghino, J.L. and Gold, M.H. 1999. 1,4-benzoquinone reductase from *Phanerochaete chry-sosporium*: cDNA cloning and regulation of expression. Appl. Environ. Microb. 65: 415–421. Altschul, S.F., Madden, T.L., Schaffer, A.A., Zhang, J., Zhang, Z., Miller, W. and Lipman, D.J. 1997. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25: 3389–3402.

- Boerjan, W., Ralph, J. and Baucher, M. 2003. Lignin biosynthesis. Annu. Rev. Plant Biol. 54: 519–546.
- Bruce, W., Desbons, P., Crasta, O. and Folkerts, O. 2001. Gene expression profiling of two related maize inbred lines with contrasting root-lodging traits. J. Exp Bot. 52: 459–468.
- Chang, W.W., Huang, L., Shen, M., Webster, C., Burlingame, A.L. and Roberts, J.K. 2000. Patterns of protein synthesis and tolerance of anoxia in root tips of maize seedlings acclimated to a low-oxygen environment, and identification of proteins by mass spectrometry. Plant Physiol. 122: 295–318.
- Clauser, K.R., Baker, P.R. and Burlingame, A.L. 1999. Role of accurate mass measurement (\pm 10 ppm) in protein identification strategies employing MS or MS/MS and database searching. Anal. Chem. 71: 2871–2882.
- Damerval, C., Vienne, D., Zivy, M. and Thiellement, H. 1986. Technical improvements in two-dimensional electrophoresis increase the level of genetic variation detected in wheat seedling proteins. Electrophoresis 7: 52–54.
- Esau, K. 1965. Plant Anatomy. John Wiley and Sons, New York.
- Goddemeier, M.L., Wulff, D. and Feix, G. 1998. Root specific expression of a *Zea mays* gene encoding a novel glycine-rich protein, ZmGrp3. Plant Mol. Biol. 36: 799–802.
- Hetz, W., Hochholdinger, F., Schwall, M. and Feix, G. 1996. Isolation and characterisation of *rtcs*, a mutant deficient in the formation of nodal roots. Plant J. 10: 845–857.
- Hochholdinger, F. and Feix, G. 1998. Early post-embryonic root formation is specifically affected in the maize mutant *lrt1*. Plant J. 16: 247–255.
- Hochholdinger, F., Park, W.J., Sauer, M. and Woll, K. 2004a. From weeds to crops: genetic analysis of root development in cereals. Trends Plant Sci. 9: 42–48.
- Hochholdinger, F., Guo, L. and Schnable, P.S. 2004b. Cytoplasmic regulation of the accumulation of nuclear-encoded proteins in the mitochondrial proteome of maize. Plant J. 37: 199–208.
- Hose, E., Clarkson, D.T., Steudle, E., Schreiber, L. and Hartung, W. 2001. The exodermis: a variable apoplastic barrier. J. Exp. Bot. 52: 2245–2264.
- Jung, E., Heller, M., Sanchez, J.C. and Hochstrasser, D.F. 2000. Proteomics meets cell biology: the establishment of subcellular proteomes. Electrophoresis 21: 3369–3377.

- Lynch, J. 1995. Root architecture and plant productivity. Plant Physiol. 109: 7–13.
- Matsuyama, T., Yasumura, N., Funakoshi, M., Yamada, Y. and Hashimoto, T. 1999a. Maize genes specifically expressed in the outermost cells of root cap. Plant Cell Physiol. 40: 469–476.
- Matsuyama, T., Satoh, H., Yamada, Y. and Hashimoto, T. 1999b. A maize glycine-rich protein is synthesized in the lateral root cap and accumulates in the mucilage. Plant Physiol. 120: 665–674.
- McCully, M.E. and Canny, M.J. 1988. Pathways and processes of water and nutrient movements in roots. Plant Soil 111: 159–170.
- Ponce, G., Luján, R., Campos, M.E., Reyes, A., Nieto-Sotelo, J., Feldman, L.J. and Cassab, G.I. 2000. Three maize rootspecific genes are not correctly expressed in regenerated caps in the absence of the quiescent center. Planta 211: 23–33.
- Porubleva, L., Van der Velden, K., Kothari, S., Oliver, D.J. and Chitnis, P.R. 2001. The proteome of maize leaves: use of gene sequences and expressed sequence tag data for identification of proteins with peptide mass fingerprints. Electrophoresis 22: 1724–1738.
- Reed, R.C., Brady, S.R. and Muday, G.K. 1998. Inhibition of auxin movement from the shoot into the root inhibits lateral root development in *Arabidopsis*. Plant Physiol. 118: 1369–1378.
- Schoof, H., Zaccaria, P., Gundlach, H., Lemcke, K., Rudd, S., Kolesov, G., Arnold, R., Mewes, H.W. and Mayer, K.F. 2002. MIPS *Arabidopsis thaliana* Database (MAtDB): an integrated biological knowledge resource based on the first complete plant genome. Nucleic Acids Res. 30: 91–93.
- Varney, G.T. and Canny, M.J. 1993. Rates of water uptake into the mature root system of maize plants. New Phytol. 123: 775–786.
- Wang, X.L., Canny, M.J. and McCully, M.E. 1991. The water status of the roots of soil-grown maize in relation to the maturity of their xylem. Physiol. Plant 82: 157–162.
- Wang, X.L., McCully, M.E. and Canny, M.J. 1994. The branch roots of Zea. IV. The maturation and openness of xylem conduits in first-order branches of soil-grown roots. New Phytol. 126: 21–29.
- Wang, X.L., McCully, M.E. and Canny, M.J. 1995. Branch Roots of Zea. V. Structural features that may influence water and nutrient transport. Bot. Acta 108: 209–219.